FORM PTO-1390 U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARS (REV. 4-2001)	OFFICE ATTORNEY 'S DOCKET NUMBER								
TRANSMITTAL LETTER TO THE UNITE	D STATES 59138-B-PCT-US/JPW/FHB								
DESIGNATED/ELECTED OFFICE (DO									
CONCERNING A FILING UNDER 35 U.									
INTERNATIONAL APPLICATION NO. INTERNATIONAL FI									
PCT/US00/12065 3 May 2000	3 May 1999								
TITLE OF INVENTION DNA ENCODING SNORF36a AND SNORF 36b RECEPTORS									
APPLICANT(S) FOR DO/EO/US SYNAPTIC PHARMACEUTICAL CORPORATION									
Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information:									
1. This is a FIRST submission of items concerning a filing under 35 U.S.C. 371.									
2. This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U.S.C. 371.									
3. This is an express request to begin national examination proce items (5), (6), (9) and (21) indicated below.	This is an express request to begin national examination procedures (35 U.S.C. 371(f)). The submission must include items (5) (6), (9) and (21) indicated below.								
4. The US has been elected by the expiration of 19 months from									
5. A copy of the International Application as filed (35 U.S.C. 37)									
. 57 1 1	d by the International Bureau). (courtesy copy)								
b. X has been communicated by the International Bureau. c. is not required, as the application was filed in the Un									
Communication of the American American									
a. is attached hereto.									
b. has been previously submitted under 35 U.S.C. 154(d)(4).								
7. Amendments to the claims of the International Aplication under									
a. are attached hereto (required only if not communicate	ed by the International Bureau).								
b. have been communicated by the International Bureau	ì.								
c. have not been made; however, the time limit for mak	ing such amendments has NOT expired.								
d. X have not been made and will not be made.									
8. An English language translation of the amendments to the claim	as under PCT Article 19 (35 U.S.C. 371 (c)(3)).								
9. An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)).	(unsigned)								
10. An English lanugage translation of the annexes of the Internation Article 36 (35 U.S.C. 371(c)(5)).	O. An English lanugage translation of the annexes of the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(c)(5)).								
Items 11 to 20 below concern document(s) or information includ	ed:								
11. An Information Disclosure Statement under 37 CFR 1.97 and	1.98.								
12. An assignment document for recording. A separate cover she	et in compliance with 37 CFR 3.28 and 3.31 is included.								
13. A FIRST preliminary amendment.									
14. A SECOND or SUBSEQUENT preliminary amendment.									
15. A substitute specification.									
16. A change of power of attorney and/or address letter.	A change of power of attorney and/or address letter.								
17. A computer-readable form of the sequence listing in accordan	A computer-readable form of the sequence listing in accordance with PCT Rule 13ter.2 and 35 U.S.C. 1.821 - 1.825.								
A second copy of the published international application under 35 U.S.C. 154(d)(4).									
Other items of information: Courtesy copy of Notice Informing Applicant of the Communication of the International Application to the Designated Offices, PCT Request, PCT Demand, International reliminary Examination Report, Renewed Petition Under 37 C.F.R. 1.181 and Petition For Three Month extension of Time Under 37 C.F.R. 1.136(a), Statement in Accordance with 37 C.F.R. 1.821(f), and mall Entity Declaration from U.S. parent application, 1 loose set of formal drawings, and an express Mail Certificate of Mailing bearing Label No. EK 873 630 962 US dated November 1, 2001									

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U.S. APPLICATION NO. (If known). Sec 37 CER 1.5) INTERNATIONAL APPLICATION NO. not yet known 8 9 2 PCT/US00/12065					ATTORNEY'S DOCKET NUMBER 59138-B-PCT-US/JPW				
	21. X The following fees are submitted:					PTO USE ONLY	7		
	BASIC NATIONAL	L FEE (37 CFR 1.	492 (a) (1) - (5)):						
	Neither internation	nal preliminary exa							
nor international search fee (37 CFR 1.445(a)(2)) paid to USPTO and International Search Report not prepared by the EPO or JPO\$1040.00									
International preliminary examination fee (37 CFR 1.482) not paid to USPTO but International Search Report prepared by the EPO or JPO\$890.00									
International preliminary examination fee (37 CFR 1.482) not paid to USPTO but international search fee (37 CFR 1.445(a)(2)) paid to USPTO									
International preliminary examination fee (37 CFR 1.482) paid to USPTO but all claims did not satisfy provisions of PCT Article 33(1)-(4) \$710.00									
International preliminary examination fee (37 CFR 1.482) paid to USPTO and all claims satisfied provisions of PCT Article 33(1)-(4)\$100.00									
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ł	CLAIMS Total claims	18 - 20		x \$18.00	\$	0.00		_	
ł	Independent claims	3 - 3 =		x \$84.00	\$	0.00			
	MULTIPLE DEPENI	DENT CLAIM(S)	if applicable)	+ \$280.00	\$	0.00			
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Applicant claims small entity status. See 37 CFR 1.27. The fees indicated above are reduced by 1/2.					\$ 3.	55.00			
t				JBTOTAL =	\$ 3.	55.00	<u> </u>	_	
Processing fee of \$130.00 for furnishing the English translation later than 20 30 months from the earliest claimed priority date (37 CFR 1.492(f)).					\$	0.00			
TOTAL NATIONAL FEE =					\$	0.00			
Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31). \$40.00 per property +					\$	0.00	1		
TOTAL FEES ENCLOSED =				\$ 3.	55.00				
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						charged:	\$		
a. X A check in the amount of \$ 355.00 to cover the above fees is enclosed.									
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b. Please charge my Deposit Account No in the amount of \$ to cover the above fees. A duplicate copy of this sheet is enclosed.									
c. X The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 03-3125. A duplicate copy of this sheet is enclosed.									
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	NUIE: Where an a 1.137 (a) or (b)) mus	t be filed and grai	nt under 37 CFR 1.494 of 1 nted to restore the application	n to pending status.	\\\	\ \	L		
SEND ALL CORRESPONDENCE TO:									
John P. White, Esq. SIGNATUR									
Reg. No. 28,678					pl. w	hite			
Cooper & Dunham LLP NAME									
1185 Avenue of the Americas New York 10036 28,67						'8			
New York, New York 10036 REGISTRATION NUMBER									
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Dkt. 59138-B-PCT-US/JPW/FHB

IN THE UNITED STATES DESIGNATED ELECTED OFFICE (DO/EO/US)

Applicants

Beth E. Borowsky, et al.

International

Application No. :

PCT/US00/12065

Filed

3 May 2000

For

DNA ENCODING SNORF36a AND SNORF 36b

RECEPTORS

1185 Avenue Of The Americas New York, New York 10036

November 1, 2001

Assistant Commissioner for Patents

Washington, D.C. 20231

Attn: DO/EO/US

Sir:

PRELIMINARY AMENDMENT

Applicants request that the following amendments be made in the above-identified application:

In the Claims:

Please cancel claims 15-18, 20-35, 37-46, 48-57, and 59-165 without disclaimer or prejudice to applicants' right to pursue the subject matter of these claims in a future continuation or divisional application.

Beth E. Borowsky, et al. U.S. Serial No.: Not Yet Known (§371 National Stage Application of PCT/US00/12065, filed 3 May 2000) Filed: Herewith Page 2

REMARKS

This application is a §371 national stage of PCT International Application No. PCT/US00/12065, filed 3 May 2000, designating the United States of America, which claims priority of U.S. Serial Nos. 09/518,914, filed March 3, 2000, and 09/303,593, filed May 3, 1999. Accordingly, the parent application, PCT International Application No. PCT/US00/12065, is pending today in the United States of America pursuant to 35 U.S.C. §363, and the subject §371 national stage application is co-pending therewith in fulfillment of the provisions of 35 U.S.C. §120.

Sequence Listing

The Sequence Listing in the subject application is identical to the Sequence Listing in the parent of the subject application, namely PCT International Application No. PCT/US00/12065, filed May 3, 2000. Applicants attach herewith copies of the paper copy of the Sequence Listing (20 pages) and the Statement In Accordance With 37 C.F.R. §1.821(f) which were filed with PCT International Application No. PCT/US00/12065 on May 3, 2000. In accordance with 37 C.F.R. §1.821 (e), please use the computer readable form filed in PCT International Application No. PCT/US00/12065 on May 3, 2000 as the computer readable form for the instant application. It is understood that the Patent and Trademark Office will make the necessary change in application number and filing date for the computer readable form that will be used for the instant application.

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Beth E. Borowsky, et al. U.S. Serial No.: Not Yet Known (§371 National Stage Application of PCT/US00/12065, filed 3 May 2000) Filed: Herewith Page 3

Claims 1-165 were pending in the subject application. By this Amendment, applicants have canceled claims 15-18, 20-35, 37-46, 48-57, and 59-165 without prejudice or disclaimer. Accordingly, upon entry of this Amendment, claims 1-14, 19, 36, 47, 58 will be pending and under examination.

If a telephone interview would be of assistance in advancing prosecution of the subject application, applicants' undersigned attorney invites the Examiner to telephone him at the number provided below.

No fee, other than the enclosed \$355.00 filing fee for the subject application, is deemed necessary in connection with filing this Preliminary Amendment. However, if an additional fee is required, authorization is hereby given to charge the amount of any such fee to Deposit Account No. 03-3125.

Respectfully submitted,

John /P. White

Registration No. 28,678 Attorney for Applicants

Cooper & Dunham LLP

1185 Avenue of the Americas New York, New York 10036

(212) 278-0400

10048192**10/0181**92

JC13 Rec'd PCT/PTO 0 1 NOV 2001

1795/59138-B-PCT-US/JPW/FHB

Application for United States Tetters Patent

To all whom it may concern:

Be it known that

Beth E. Borowsky, et al.

have invented certain new and useful improvements in

DNA ENCODING SNORF36a AND SNORF36b RECEPTORS

of which the following is a full, clear and exact description.

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10/018192 PC1/US00/12065

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JC13 Rec'd PCT/PTO 0 1 NOV 2001

DNA ENCODING SNORF36a AND SNORF36b RECEPTORS

BACKGROUND OF THE INVENTION

This application is a continuation-in-part and claims priority of U.S. Serial No. 09/518,914, filed March 3, 2000, a continuation-in-part of U.S. Serial No. 09/303,593, filed May 3, 1999, the contents of which are hereby incorporated by reference into the subject application.

Throughout this application various publications are referred to by partial citations within parenthesis. Full citations for these publications may be found at the end of the specification immediately preceding the claims. The disclosures of these publications, in their entireties, are hereby incorporated by reference into this application in order to more fully describe the state of the art to which the invention pertains.

Neuroregulators comprise a diverse group of natural products that subserve or modulate communication in the nervous system. They include, but are not limited to, neuropeptides, amino acids, biogenic amines, lipids and lipid metabolites, and other metabolic byproducts. Many of these neuroregulator substances interact with specific cell surface receptors which transduce signals from the outside to the inside of the cell. G-protein coupled receptors (GPCRs) represent a major of cell surface receptors with which class neurotransmitters interact to mediate their effects. GPCRs are characterized by seven membrane-spanning domains and are coupled to their effectors via G-proteins linking receptor activation with intracellular biochemical sequelae such as stimulation of adenylyl cyclase.

Opsins represent one of the major families of GPCRs. These

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receptors are unique compared to other GPCRs in that light crucial co-factor for their activation physiological conditions. A major subclass of the opsin family is that of visual opsins such as rhodopsin and cone visual opsins, also known as The photopigments, are located in the eye and are involved in transducing visual information from the eye to the brain. Our understanding of opsin function has been derived primarily from the study of visual photopigments.

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Rhodopsin and cone opsins are localized in retinal rod and cone photoreceptors, respectively. These photopigments respond to different wavelengths of light and thus have very distinct absorption spectra associated with different absorption maxima (λ_{max}) . Even though both receptor subtypes convey visual signals to the brain in response illumination, they have evolved to perform very distinct functions related to vision. Cone opsins are primarily responsible for color vision, also known as photopic vision, in different species. In contrast, rhodopsin, believed to have evolved from cone opsin, is mainly involved in dim light vision, also known as scotopic vision. Rhodopsin, highly enriched in rod photoreceptor membranes, has been used extensively as a model receptor to understand activation mechanism and functioning of opsins.

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Rhodopsin contains the seven membrane-spanning apoprotein opsin and a retinoid-based chromophore (See reviews Hargrave and McDowell, 1992; Yarfitz and Hurley, 1994). In the ground or inactive state (i.e. in the absence of light), the chromophore, usually 11-cis-retinal, is covalently attached to a highly conserved lysine residue in the middle of the seventh transmembrane segment via a protonated Schiff base. All vertebrate visual opsins contain a highly conserved glutamate residue in the transmembrane helix 3 which serves

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as a counterion for the protonated Schiff base. It has been postulated that 11-cis-retinal behaves as an inverse agonist and induces an inactive conformation of the apoprotein which, by itself, is partially active (Cohen et al., 1993; Surya et Upon absorbing a photon, 11-cis-retinal is al., 1995). isomerized to the agonist all-trans-retinal which introduces distortion in the opsin and initiates a cascade conformational changes in the molecule. Rhodopsin is first converted to bathorhodopsin, followed by lumirhodopsin, metarhodopsin I and metarhodopsin II states in a sequential manner. Even though most of these transient conformational states are difficult to study biochemically, they can be easily distinguished on the basis of their spectroscopic properties since each state has a unique absorption maximum. Experimental evidence suggests that the formation metarhodopsin II, a relatively stable state, deprotonation of the Schiff base and represents the active conformation of the apoprotein. In this state, the opsin the cognate G-protein and activates initiates intracellular signaling cascade which ultimately results in transfer of visual information to the brain. Upon hydrolysis of the Schiff base linkage, metarhodopsin II decays into free all-trans-retinal and opsin. All-trans-retinal transported to the neighbouring retinal pigment epithelial cells where it is converted to 11-cis-retinal via enzymatic 11-cis-retinal is transported back to retinal photoreceptors where it recombines with the opsin apoprotein to regenerate the rhodopsin molecule.

30 Even though all visual opsins essentially use the same activation mechanism as rhodopsin, there are some noticeable differences between vertebrate and invertebrate visual opsins (Gartner and Towner, 1995; Yarfitz and Hurley, 1994, Terakita et al., 1998; Arnheiter, 1998). Activation of vertebrate visual pigment results primarily in stimulation of G_t G-

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protein (also known as transducin) leading to an increase in phosphodiesterase activity. Initiation of signaling cascade ultimately results in closure of cation channels and hyperpolarization of the cell. In contrast, opsin visual pigments in invertebrates such as squid and fruitfly activate Ga G-protein and elevate intracellular IP3 and Ca²⁺ levels (Wood et al., 1989; Nobes et al., 1992; Yarfitz and Hurley, 1994). Another major difference between vertebrate and invertebrate visual opsins is the stability of the active conformation of the receptor. Formation of vertebrate metarhodopsin II, the active conformation of rhodopsin, is rapidly followed by hydrolysis of the Schiff base linkage and dissociation of metarhodopsin II into free all-trans-retinal and opsin apoprotein. It has been suggested that the glutamate counterion in the transmembrane helix 3 aids in the hydrolysis reaction (Gartner and Towner, In contrast, invertebrate metarhodopsin represents a thermally stable state where the chromophore remains attached to the apoprotein (Kiselev and Subramaniam, 1994). This allows rapid photoisomerization of all-trans-retinal back to 11-cis-retinal within the apoprotein and rapid regeneration of rhodopsin, thus eliminating the need for retinal regenerating tissue (Provencio et al., 1998). thermally stable metastate of invertebrate photopigment may be formed due to the absence of the glutamate counterion in transmembrane helix 3 of invertebrate visual opsins (Gartner and Towner, 1995).

Most opsins use 11-cis-retinal derived from carotenoids as a chromophore; however, some opsins use 3-hydroxy, 4-hydroxy or 3, 4-dehydro isomers of 11-cis-retinal as a chromophore to accommodate the abundant availability of the substituted carotenoids (Gartner and Towner, 1995). Different opsins respond to photons with different wavelengths, a phenomenon known as spectral tuning. Even though the use of a

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particular retinal derivative as a chromophore contributes to spectral specificity (Gartner and Towner, 1995), the major determinant of spectral tuning is the presence of unique acids surrounding the retinal-binding (Kochendoerfer et al., 1999). For example, substitution of a highly conserved glycine in transmembrane helix 3 of rhodopsin with amino acids of increasing size results in progressive shift of λ_{max} towards the blue wavelength (Han et al., 1996). Similarly, replacement of conserved non-polar residues with hydroxyl amino acids changes the opsin from a green-absorbing molecule to a red-absorbing pigment (Chan et al., 1992).

Even though the visual opsins have been at the forefront of opsin research, scientists are now turning their attention to non-visual opsins (the opsins not involved in transducing visual information) because of their potential involvement in physiological processes such as circadian rhythm and The existence of non-visual photopigments in reproduction. nonmammalian vertebrates was first suggested by Karl von Frisch. He demonstrated that the skin of the European minnow changed color in response to light even in the absence of the eye and pineal gland, and postulated photoreceptive elements in the diencephalon (Foster et al, 1994). Further evidence supporting the presence of non-visual photopigments was obtained in blinded lampreys and ducks which responded to illumination with body movements and gonadal induction, respectively (Foster et al., 1994). Recent studies using histochemical techniques has further corroborated these physiological observations. Silver et al. immunostained the cerebrospinal fluid (CSF)-contacting neurons with anti-opsin antibody in brains of the ring dove, quail and duck. Similarly, intense immunostaining of the CSF-contacting neurons within the basal brain of the lizard, Anolis carolinensis, was observed with anti-cone opsin

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antibody (Foster et al., 1993).

Recent molecular cloning of several non-visual opsins is in agreement with the above-mentioned studies. Pinopsin is expressed in the pineal gland of the chicken and is believed to play a role in circadian rhythm (Okano et al., 1994; Max et al., 1995). Interestingly, expression of the pinopsin gene is regulated by light (Takanaka et al., 1998). Max et al. (1998) have demonstrated light-dependent activation of transducin by pinopsin, implying that the pinopsin is a functional photoreceptor. Two other opsins identified in pinealocytes are vertebrate ancient (VA) opsin cloned from the salmon fish (Soni et al., 1997; Soni et al., 1998), and parapinopsin cloned from the channel catfish (Blackshaw and Snyder, 1997). In addition to pineal cells, VA opsin is also localized in the amacrine and horizontal cells of the salmon retina. On the other hand, expression of parapinopsin is confined to the parapineal and pineal organs of the catfish.

Several of the non-visual opsins are, in fact, expressed in the eye. Sun et al. (1997) cloned peropsin from human retina and mouse eye. This opsin is localized exclusively in microvilli of the apical membrane of retinal pigment epithelial (RPE) cells, indicating that it may function as a sensor of retinoids generated in the adjacent outer membrane of rhodopsin or cone opsins. RPE retinal G-protein-coupled receptor (RGR) is another receptor found in the RPE (Tao et al., 1998). Unlike other opsins which are believed to be present at the plasma membrane, RGR is localized intracellularly. The amino acid sequence of RGR suggests that, along with squid retinochrome, it may form a distinct subfamily of opsins (Hara-Nishimura et al., 1990).

Interestingly, it has been suggested that RGR may prefer alltrans-retinal, rather than 11-cis-retinal, as a ligand and

-7-

may be involved in the photoisomerization of the all-trans isomer to the 11-cis isomer (Hao and Fong, 1999). In such a case, its function may be the rapid regeneration of 11-cis-retinal in the RPE for use in the visual cycle.

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One of the known photo-sensitive processes is melanosome dispersion in the dermal melanophores of Xenopus laevis. accordance with this, melanopsin has been cloned from melanophores (Provencio et al., 1998). Melanopsin is expressed in the melanophores, suprachiasmatic and preoptic nuclei of the hypothalamus, iris, RPE and retina. expression in visual and nonvisual tissues suggests a role in visual and nonvisual photosensory phenomena. a non-visual opsin has been identified in the mammalian Blackshaw and Snyder (1999)encephalopsin, which, as the name suggests, is highly expressed in various areas of the brain. It is present in the preoptic area and the paraventricular nucleus of the hypothalamus, the cerebral cortex, cerebellar Purkinje cells, striatum, thalamus and the ventral horn of the spinal cord. Interestingly, this receptor is not present in the eye.

The molecular identification of non-visual opsins has raised several questions. How are they activated? What is their physiological function? All the non-visual opsins cloned to date contain lysine in the seventh transmembrane helix, the site for retinal chromophore attachment, implying that a retinoid may be the chromophore for the non-visual opsins, similar to the visual opsins. Several groups have been successful in reconstituting non-visual opsins with retinoids and activating them with light (Okano et al., 1994; Soni et al., 1998). Retinoids can cross the blood-brain barrier. albeit at low efficiency (Pardridge et al., 1985; Franke et Furthermore, a transporter with high affinity al., 1999). for retinoids, β-Trace, has been recognized recently (Tanaka

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et al., 1997). This secretory protein is present in high levels in the CSF and may transport retinoids to different regions of the brain, analogous to the plasma RBP. That retinoids are indeed present in the brain was demonstrated by Foster et al. (1993) who were able to identify retinal isomers in the Anolis anterior brain using HPLC analysis. If a retinal isomer is indeed a chromophore for non-visual opsins then light would be needed to photoisomerize the isomer and activate the receptor. Several reports suggest that light can reach the deep areas of the brain (Muller and Wilson, 1986; Grace et al., 1996; Blackshaw and Snyder, 1999), and a neurotransmitter release-enhancing effect of light on cortical slices has been observed (Wade et al., Therefore, the activation mechanisms of non-visual opsins may be similar to the visual opsins. However, it should be noted here that some non-visual opsins have proven resistant to functional reconstitution with retinal isomers (Provencio et al., 1998; Blackshaw and Snyder, 1999), raising the possibility that these receptors utilize a non-retinoid ligand and may not require light for activation.

What could be the function of non-visual opsins? interesting possibility is that they may be involved in Circadian rhythm represents circadian rhythm. fluctuations in biological activities that are regulated by the light-dark cycles. It is composed of three components: a photoactive input, the circadian clock itself which exhibits periodicity, and the behavioral and physiological oscillations as output. In animals, in the phenomenon known as photoentrainment, exposure to light results in regulation However, the identity of the of the circadian rhythm. photoreceptive molecule mediating photoentrainment Since the photoentrainment response remained a mystery. occurs at the wavelength of 500 nm, it has been suggested that an opsin may be mediating the response (Foster, 1998).

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Freedman et al. (1999) and Lucas et al. (1999) recently demonstrated that photoentrainment was intact in mice lacking rod and cone receptors; however, removal of the eyes in these mice abolished the effect of light on circadian rhythm as well as on melatonin synthesis. These results, and ocular localization of several non-visual opsins, strongly support the role of ocular non-visual opsins in photoentraining circadian rhythm. In addition, the non-visual opsins localized in the CNS may form a component of the circadian clock itself.

If non-visual opsins are indeed involved in regulating circadian rhythm, then they represent an attractive therapeutic target for circadian rhythm-related conditions. These include sleep disorders such as jet lag. It has been suggested that a change in circadian rhythm may be an underlying cause for sleep disorders such as insomnia, Advanced Sleep Phase Syndrome and Delayed Sleep Phase Syndrome (Sedgwick, 1998; Refinetti, 1999). In addition, dissociation between biological clock and work hours may result in shift-work-related sleep disorders. Importantly, bright light therapy has been demonstrated to help in these disorders (Rosenthal et al., 1990; Lack and Wright, 1993; Campbell et al., 1995; Murphy and Campbell, 1996; Cooke et al., 1998; Refinetti, 1999). Similarly, exposure to light at appropriate time reduces the effect of jet lag on travelers (Refinetti, 1999). These observations suggest that non-visual opsins may mediate these beneficial effects of light in circadian rhythm-related disorders.

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Non-visual opsins also may play a role in seasonal affective disorder (SAD). This disorder is characterized by a subpopulation of people suffering from depression during winter. Light therapy is effective in these people (Terman and Terman, 1999); especially green light is more effective than red light (Oren et al., 1991). It has been recently

-10-

hypothesized that the interaction of specialized photoreceptors with magnetic field may influence sensitivity of patients suffering from SAD to light (Partonen, 1998).

In addition, the discreet localization of various opsins in the CNS areas indicate their potential role in CNS-related physiology and disorders.

Non-visual photoreceptors are also involved in melanosome dispersion in melanophores, and thus, in change in the color of the skin in various species. In birds and mammals, non-visual 'deep brain photoreceptors' are also linked to reproductive behaviour and photoperiodic gonadal responses (Yoshikawa and Oishi, 1998).

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In summary, opsins constitute an important branch of the GPCR superfamily. They behave as photosensitive elements. They are localized in the retina and in non-retinal locations including the brain. The retinal rod and cone opsins are mainly responsible for conveying visual information to the brain, while the non-visual opsins in the retina and elsewhere may be involved in regulation of melatonin synthesis and circadian rhythm, photoentrainment, SAD, skin colour change and camouflage, and reproductive behaviour.

-11-

SUMMARY OF THE INVENTION

This invention provides an isolated nucleic acid encoding a mammalian SNORF36 receptor.

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This invention further provides a purified mammalian SNORF36 receptor protein.

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This invention also provides a vector comprising a nucleic acid in accordance with this invention.

This invention still further provides a cell comprising a vector in accordance with this invention.

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This invention additionally provides a membrane preparation isolated from a cell in accordance with this invention.

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Furthermore, this invention provides a nucleic acid probe comprising at least 15 nucleotides, which probe specifically hybridizes with a nucleic acid encoding a mammalian SNORF36 receptor, wherein the probe has a sequence complementary to a unique sequence present within one of the two strands of the nucleic acid encoding the mammalian SNORF36 receptor contained in plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. 203977), plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. 203976), plasmid pEXJ.T3T7-rSNORF36p (ATCC Patent Depository No. PTA-534) or plasmid pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216).

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This invention further provides a nucleic acid probe comprising at least 15 nucleotides, which probe specifically hybridizes with a nucleic acid encoding a mammalian SNORF36 receptor, wherein the probe has a sequence complementary to a unique sequence present within (a) the nucleic acid sequence shown in Figures 1A-1C (SEQ ID NO: 1), (b) the

-12-

reverse complement thereof.

This invention further provides a nucleic acid probe comprising at least 15 nucleotides, which probe specifically hybridizes with a nucleic acid encoding a mammalian SNORF36 receptor, wherein the probe has a sequence complementary to a unique sequence present within (a) the nucleic acid sequence shown in Figures 3A-3C (SEQ ID NO: 3), (b) the reverse complement thereof.

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This invention provides an antisense oligonucleotide having a sequence capable of specifically hybridizing to RNA encoding a mammalian SNORF36 receptor, so as to prevent translation of such RNA.

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This invention further provides an antisense oligonucleotide having a sequence capable of specifically hybridizing to genomic DNA encoding a mammalian SNORF36 receptor, so as to prevent transcription of such genomic DNA.

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This invention also provides an antibody capable of binding to a mammalian SNORF36 receptor encoded by a nucleic acid in accordance with this invention.

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Moreover, this invention provides an agent capable of competitively inhibiting the binding of an antibody in accordance with this invention to a mammalian SNORF36 receptor.

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This invention yet further provides a pharmaceutical composition comprising (a) an amount of an oligonucleotide in accordance with this invention capable of passing through a cell membrane and effective to reduce expression of a mammalian SNORF36 receptor and (b) a pharmaceutically acceptable carrier capable of passing through the cell membrane.

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-13-

This invention also provides a pharmaceutical composition which comprises an amount of an antibody in accordance with this invention effective to block binding of a ligand to a human SNORF36 receptor and a pharmaceutically acceptable carrier.

This invention further provides a transgenic, nonhuman mammal expressing DNA encoding a mammalian SNORF36 receptor in accordance with this invention.

This invention still further provides a transgenic, nonhuman mammal comprising a homologous recombination knockout of a native mammalian SNORF36 receptor.

This invention further provides a transgenic, nonhuman mammal whose genome comprises antisense DNA complementary to DNA encoding a mammalian SNORF36 receptor in accordance with this invention so placed within such genome as to be transcribed into antisense mRNA which is complementary to and hybridizes with mRNA encoding the mammalian SNORF36 receptor so as to reduce translation of such mRNA and expression of such receptor.

This invention provides a process for identifying a chemical compound which specifically binds to a mammalian SNORF36 receptor which comprises contacting cells containing DNA encoding, and expressing on their cell surface, the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with the compound under conditions suitable for binding, and detecting specific binding of the chemical compound to the mammalian SNORF36 receptor.

This invention further provides a process for identifying a chemical compound which specifically binds to a mammalian SNORF36 receptor which comprises contacting a membrane

-14-

preparation from cells containing DNA encoding, and expressing on their cell surface, the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with the compound under conditions suitable for binding, and detecting specific binding of the chemical compound to the mammalian SNORF36 receptor.

This invention still further provides a process involving competitive binding for identifying a chemical compound which specifically binds to a mammalian SNORF36 receptor which comprises separately contacting cells expressing on their cell surface the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with both the chemical compound and a second chemical compound known to bind to the receptor, and with only the second chemical compound, under conditions suitable for binding of such compounds to the receptor, and detecting specific binding of the chemical compound to the mammalian SNORF36 receptor, a decrease in the binding of the second chemical compound to the mammalian SNORF36 receptor in the presence of the chemical compound being tested indicating that such chemical compound binds to the mammalian SNORF36 receptor.

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invention further provides a process involving competitive binding for identifying a chemical compound which specifically binds to a mammalian SNORF36 receptor which comprises separately contacting a membrane preparation from cells expressing on their cell surface the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with both the chemical compound and a second chemical compound known to bind to the receptor, and with only the second chemical compound, under conditions suitable for binding of such compounds to the receptor, and detecting specific binding of the chemical compound to the

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-15-

mammalian SNORF36 receptor, a decrease in the binding of the second chemical compound to the mammalian SNORF36 receptor in the presence of the chemical compound being tested indicating that such chemical compound binds to the mammalian SNORF36 receptor.

This invention further provides a compound identified by one of the processes of this invention.

This invention provides a method of screening a plurality of chemical compounds not known to bind to a mammalian SNORF36 receptor to identify a compound which specifically binds to the mammalian SNORF36 receptor, which comprises (a) contacting cells transfected with, and expressing, DNA encoding the mammalian SNORF36 receptor with a compound known to bind specifically to the mammalian SNORF36 receptor; (b) contacting the cells of step (a) with the plurality of compounds not known to bind specifically to the mammalian SNORF36 receptor, under conditions permitting binding of compounds known to bind to the mammalian SNORF36 receptor; (c) determining whether the binding of the compound known to bind to the mammalian SNORF36 receptor is reduced in the presence of the plurality of compounds, relative to the binding of the compound in the absence of the plurality of compounds; and if so (d) separately determining the binding to the mammalian SNORF36 receptor of each compound included in the plurality of compounds, so as to thereby identify any compound included therein which specifically binds to the mammalian SNORF36 receptor.

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This invention further provides a method of screening a plurality of chemical compounds not known to bind to a mammalian SNORF36 receptor to identify a compound which specifically binds to the mammalian SNORF36 receptor, which comprises (a) contacting a membrane preparation from cells transfected with, and expressing, DNA encoding the mammalian

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-16-

SNORF36 receptor with the plurality of compounds not known to bind specifically to the mammalian SNORF36 receptor under conditions permitting binding of compounds known to bind to the mammalian SNORF36 receptor; (b) determining whether the binding of a compound known to bind to the mammalian SNORF36 receptor is reduced in the presence of the plurality of compounds, relative to the binding of the compound in the absence of the plurality of compounds; and if so (c) separately determining the binding to the mammalian SNORF36 receptor of each compound included in the plurality of compounds, so as to thereby identify any compound included therein which specifically binds to the mammalian SNORF36 receptor.

This invention also provides a method of detecting expression of a mammalian SNORF36 receptor by detecting the presence of mRNA coding for the mammalian SNORF36 receptor which comprises obtaining total mRNA from the cell and contacting the mRNA so obtained with a nucleic acid probe according to this invention under hybridizing conditions, detecting the presence of mRNA hybridized to the probe, and thereby detecting the expression of the mammalian SNORF36 receptor by the cell.

This invention further provides a method of detecting the presence of a mammalian SNORF36 receptor on the surface of a cell which comprises contacting the cell with an antibody according to this invention under conditions permitting binding of the antibody to the receptor, detecting the presence of the antibody bound to the cell, and thereby detecting the presence of the mammalian SNORF36 receptor on the surface of the cell.

This invention still further provides a method of determining the physiological effects of varying levels of activity of mammalian SNORF36 receptors which comprises producing a

PCT/US00/12065

-17-

transgenic, nonhuman mammal in accordance with this invention whose levels of mammalian SNORF36 receptor activity are varied by use of an inducible promoter which regulates mammalian SNORF36 receptor expression.

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This invention additionally provides a method of determining the physiological effects of varying levels of activity of mammalian SNORF36 receptors which comprises producing a panel of transgenic, nonhuman mammals in accordance with this invention each expressing a different amount of mammalian SNORF36 receptor.

Moreover, this invention provides a method for identifying an antagonist capable of alleviating an abnormality wherein the abnormality is alleviated by decreasing the activity of a mammalian SNORF36 receptor comprising administering a compound to a transgenic, nonhuman mammal according to this invention, and determining whether the compound alleviates any physiological and/or behavioral abnormality displayed by the transgenic, nonhuman mammal as a result of overactivity of a mammalian SNORF36 receptor, the alleviation of such an abnormality identifying the compound as an antagonist.

This invention also provides an antagonist identified by the preceding method.

This invention further provides a composition, e.g. a pharmaceutical composition, comprising an antagonist according to this invention and a carrier, e.g. a pharmaceutically acceptable carrier.

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This invention additionally provides a method of treating an abnormality in a subject wherein the abnormality is alleviated by decreasing the activity of a mammalian SNORF36 receptor which comprises administering to the subject an effective amount of the pharmaceutical composition according

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PCT/US00/12065

-18-

to this invention so as to thereby treat the abnormality.

In addition, this invention provides a method for identifying an agonist capable of alleviating an abnormality in a subject wherein the abnormality is alleviated by increasing the activity of a mammalian SNORF36 receptor comprising administering a compound to a transgenic, nonhuman mammal according to this invention, and determining whether the compound alleviates any physiological and/or behavioral abnormality displayed by the transgenic, nonhuman mammal, the alleviation of such an abnormality identifying the compound as an agonist.

This invention further provides an agonist identified by the preceding method.

This invention still further provides a composition, e.g. a pharmaceutical composition, comprising an agonist according to this invention and a carrier, e.g. a pharmaceutically acceptable carrier.

Moreover, this invention provides a method of treating an abnormality in a subject wherein the abnormality is alleviated by increasing the activity of a mammalian SNORF36 receptor which comprises administering to the subject an effective amount of the pharmaceutical composition according to this invention so as to thereby treat the abnormality.

Yet further, this invention provides a method for diagnosing a predisposition to a disorder associated with the activity of a specific mammalian allele which comprises: (a) obtaining DNA of subjects suffering from the disorder; (b) performing a restriction digest of the DNA with a panel of restriction enzymes; (c) electrophoretically separating the resulting DNA fragments on a sizing gel; (d) contacting the resulting gel with a nucleic acid probe capable of

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PCT/US00/12065

-19-

specifically hybridizing with a unique sequence included within the sequence of a nucleic acid molecule encoding a mammalian SNORF36 receptor and labeled with a detectable marker; (e) detecting labeled bands which have hybridized to the DNA encoding a mammalian SNORF36 receptor to create a unique band pattern specific to the DNA of subjects suffering from the disorder; (f) repeating steps (a)-(e) with DNA obtained for diagnosis from subjects not yet suffering from the disorder; and (g) comparing the unique band pattern specific to the DNA of subjects suffering from the disorder from step (e) with the band pattern from step (f) for subjects not yet suffering from the disorder so as to determine whether the patterns are the same or different and thereby diagnose predisposition to the disorder if the patterns are the same.

This invention also provides a method of preparing a purified mammalian SNORF36 receptor according to the invention which comprises: (a) culturing cells which express the mammalian SNORF36 receptor; (b) recovering the mammalian SNORF36 receptor from the cells; and (c) purifying the mammalian SNORF36 receptor so recovered.

This invention further provides a method of preparing the purified mammalian SNORF36 receptor according to the invention which comprises: (a) inserting a nucleic acid encoding the mammalian SNORF36 receptor into a suitable expression vector; (b) introducing the resulting vector into a suitable host cell; (c) placing the resulting host cell in suitable conditions permitting the production of the mammalian SNORF36 receptor; (d) recovering the mammalian SNORF36 receptor so produced; and optionally (e) isolating and/or purifying the mammalian SNORF36 receptor so recovered.

Furthermore, this invention provides a process for determining whether a chemical compound is a mammalian

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-20-

SNORF36 receptor agonist which comprises contacting cells transfected with and expressing DNA encoding the mammalian SNORF36 receptor with the compound under conditions permitting the activation of the mammalian SNORF36 receptor, and detecting any increase in mammalian SNORF36 receptor activity, so as to thereby determine whether the compound is a mammalian SNORF36 receptor agonist.

This invention also provides a process for determining whether a chemical compound is a mammalian SNORF36 receptor antagonist which comprises contacting cells transfected with and expressing DNA encoding the mammalian SNORF36 receptor with the compound in the presence of a known mammalian SNORF36 receptor agonist, under conditions permitting the activation of the mammalian SNORF36 receptor, and detecting any decrease in mammalian SNORF36 receptor activity, so as to thereby determine whether the compound is a mammalian SNORF36 receptor antagonist.

This invention still further provides a composition, for example a pharmaceutical composition, which comprises an amount of a mammalian SNORF36 receptor agonist determined by a process according to this invention effective to increase activity of a mammalian SNORF36 receptor and a carrier, for example, a pharmaceutically acceptable carrier. In one embodiment, the mammalian SNORF36 receptor agonist is not previously known.

Also, this invention provides a composition, for example a pharmaceutical composition, which comprises an amount of a mammalian SNORF36 receptor antagonist determined by a process according to this invention effective to reduce activity of a mammalian SNORF36 receptor and a carrier, for example, a pharmaceutically acceptable carrier.

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This invention moreover provides a process for determining

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-21-

whether a chemical compound specifically binds to and activates a mammalian SNORF36 receptor, which comprises contacting cells producing a second messenger response and expressing on their cell surface the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with the chemical compound under conditions suitable for activation of the mammalian SNORF36 receptor, and measuring the second messenger response in the presence and in the absence of the chemical compound, a change, e.g. an increase, in the second messenger response in the presence of the chemical compound indicating that the compound activates the mammalian SNORF36 receptor.

invention still further provides а process determining whether a chemical compound specifically binds to and inhibits activation of a mammalian SNORF36 receptor, which comprises separately contacting cells producing a second messenger response and expressing on their cell surface the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with both the chemical compound and a second chemical compound known to activate the mammalian SNORF36 receptor, and with only the second chemical compound, under conditions suitable for activation of the mammalian SNORF36 receptor. measuring the second messenger response in the presence of only the second chemical compound and in the presence of both the second chemical compound and the chemical compound, a smaller change, e.g. increase, in the second messenger response in the presence of both the chemical compound and the second chemical compound than in the presence of only the second chemical compound indicating that the chemical compound inhibits activation of the mammalian SNORF36 receptor.

Further, this invention provides a compound determined by a process according to the invention and a composition, for

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-22-

example, a pharmaceutical composition, which comprises an amount of a mammalian SNORF36 receptor agonist determined to be such by a process according to the invention, effective to increase activity of the mammalian SNORF36 receptor and a carrier, for example, a pharmaceutically acceptable carrier.

This invention also provides a composition, for example, a pharmaceutical composition, which comprises an amount of a mammalian SNORF36 receptor antagonist determined to be such by a process according to the invention, effective to reduce activity of the mammalian SNORF36 receptor and a carrier, for example, a pharmaceutically acceptable carrier.

This invention yet further provides a method of screening a plurality of chemical compounds not known to activate a mammalian SNORF36 receptor to identify a compound which activates the mammalian SNORF36 receptor which comprises: (a) contacting cells transfected with and expressing the mammalian SNORF36 receptor with the plurality of compounds not known to activate the mammalian SNORF36 receptor, under conditions permitting activation of the mammalian SNORF36 determining whether the activity of the (b) mammalian SNORF36 receptor is increased in the presence of one or more of the compounds; and if so (c) separately determining whether the activation of the mammalian SNORF36 receptor is increased by any compound included in the plurality of compounds, so as to thereby identify each compound which activates the mammalian SNORF36 receptor.

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This invention provides a method of screening a plurality of chemical compounds not known to inhibit the activation of a mammalian SNORF36 receptor to identify a compound which inhibits the activation of the mammalian SNORF36 receptor, which comprises: (a) contacting cells transfected with and expressing the mammalian SNORF36 receptor with the plurality

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-23-

of compounds in the presence of a known mammalian SNORF36 receptor agonist, under conditions permitting activation of the mammalian SNORF36 receptor; (b) determining whether the extent or amount of activation of the mammalian SNORF36 receptor is reduced in the presence of one or more of the compounds, relative to the extent or amount of activation of the mammalian SNORF36 receptor in the absence of such one or more compounds; and if so (c) separately determining whether each such compound inhibits activation of the mammalian SNORF36 receptor for each compound included in the plurality of compounds, so as to thereby identify any compound included in such plurality of compounds which inhibits the activation of the mammalian SNORF36 receptor.

This invention also provides a composition, for example a pharmaceutical composition, comprising a compound identified by a method according to this invention in an amount effective to increase mammalian SNORF36 receptor activity and a carrier, for example, a pharmaceutically acceptable carrier.

This invention still further provides a composition, for example a pharmaceutical composition, comprising a compound identified by a method according to this invention in an amount effective to decrease mammalian SNORF36 receptor activity and a carrier, for example a pharmaceutically acceptable carrier.

Furthermore, this invention provides a method of treating an abnormality in a subject wherein the abnormality is alleviated by increasing the activity of a mammalian SNORF36 receptor which comprises administering to the subject a compound which is a mammalian SNORF36 receptor agonist in an amount effective to treat the abnormality.

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This invention additionally provides a method of treating an

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-24-

abnormality in a subject wherein the abnormality is alleviated by decreasing the activity of a mammalian SNORF36 receptor which comprises administering to the subject a compound which is a mammalian SNORF36 receptor antagonist in an amount effective to treat the abnormality.

This invention also provides a process for making a composition of matter which specifically binds to a mammalian SNORF36 receptor which comprises identifying a chemical compound using a process in accordance with this invention and then synthesizing the chemical compound or a novel structural and functional analog or homolog thereof.

This invention further provides a process for preparing a composition, for example, a pharmaceutical composition which comprises admixing a carrier, for example, a pharmaceutically acceptable carrier, and a pharmaceutically effective amount of a chemical compound identified by a process of in accordance with this invention or a novel structural and functional analog or homolog thereof.

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PCT/US00/12065

-25-

BRIEF DESCRIPTION OF THE FIGURES

Figures 1A-1C

Nucleotide sequence including sequence encoding a human SNORF36a receptor (SEQ ID NO: 1). Putative open reading frames including the shortest open reading frame are indicated by underlining two start (ATG) codons (at positions 10-12 and 67-69) and the stop codon (at positions 1444-1446). In addition, partial 5' and 3' untranslated sequences are shown.

Figures 2A-2B

Deduced amino acid sequence (SEQ ID NO: 2) of the human SNORF36a receptor encoded by the longest open reading frame indicated in the nucleotide sequence shown in Figures 1A-1C (SEQ ID NO: 1). The seven putative transmembrane (TM) regions are underlined.

Figures 3A-3C

Nucleotide sequence including sequence encoding a human SNORF36b receptor (SEQ ID NO: 3). Putative open reading frames including the shortest open reading frame are indicated by underlining two start (ATG) codons (at positions 10-12 and 67-69) and the stop codon (at positions 1477-1479). In addition, partial 5' and 3' untranslated sequences are shown.

Figures 4A-4C

Deduced amino acid sequence (SEQ ID NO: 4) of the human SNORF36b receptor encoded by the longest open reading frame indicated in the nucleotide sequence shown in Figures 3A-3C (SEQ ID NO: 3). The seven putative transmembrane (TM) regions are underlined.

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PCT/US00/12065

-26-

Figure 5

Nucleotide sequence including part of the sequence encoding a rat SNORF36 receptor (SEQ ID NO: 5).

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Deduced partial amino acid sequence (SEQ ID NO: 6) of the rat SNORF36 receptor encoded by the nucleotide sequence shown in Figure 5 (SEQ ID NO: 5). Putative transmembrane (TM) regions are underlined.

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Figures 7A-7B

Nucleotide alignment of partial sequences of human SNORF36 and rat SNORF36. Vertical lines represent identical residues.

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Figure 8

Amino acid alignment of partial sequences of human SNORF36 and rat SNORF36. Vertical lines represent identical residues and dots represent similar residues.

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Figures 9A-9C

Nucleotide sequence including sequence encoding a rat SNORF36 receptor (SEQ ID NO: 7). Putative open reading frames including the shortest open reading frame are indicated by underlining one start (ATG) codon (at positions 25-27) and the stop codon (at positions 1447-1449). In addition, partial 5' and 3' untranslated sequences are shown.

Figures 10A-10C

Deduced amino acid sequence (SEQ ID NO: 8) of the rat SNORF36 receptor encoded by the longest open reading frame indicated in the nucleotide sequence shown in Figures 9A-9C (SEQ ID NO: 7). The seven putative transmembrane (TM) regions are underlined.

PCT/US00/12065

-27-

Figures 11A-11D

Nucleotide alignment of human and rat SNORF36 receptors. Vertical lines indicate conserved residues, dots represent gaps in the alignment.

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Figures 12A-12B

Amino acid alignment of human and rat SNORF36 receptors. Vertical lines indicate conserved residues, dots represent gaps in the alignment.

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Figures 13A and 13B

Baseline Ca^{2^*} mobilization response in human SNORF36a-transfected Cos-7 cells which were either, (1) not exposed to lamp light (Figure 13A), or (2) exposed to lamp light for 90 minutes before the experiment (Figure 13B). The response was measured using Fluorometric Imaging Plate Reader (FLIPRTM). Cells were exposed to laser light at 1 second interval beginning at time 0 until the end of the trace. The first fluorescence value was subtracted from all the fluorescence values. No drug was added during the duration of the trace.

Figures 14A and 14B

Representative traces demonstrating the effect of all-trans-retinal (10µM) on intracellular Ca^{2+} in (1) human SNORF36a-transfected (Figure 14A), and (2) empty vector-transfected Cos-7 cells (Figure 14B). The response was measured using Fluorometric Imaging Plate Reader (FLIPRTM). All-trans-retinal was added at time indicated by the arrow. Baseline subtraction and negative control corrections were performed on the traces.

Figures 15A and 15B

Concentration-dependent effect of retinal analogues on intracellular Ca²⁺ in human SNORF36a-transfected (1) Cos-7

-28-

cells (n=3) (Figure 15A), and (2) HEK293 cells (n=1) (Figure 15B). Data are presented as mean \pm SD.

Figures 16A and 16B

Concentration-dependent effect of all-trans-retinal on phosphoinositide hydrolysis in human SNORF36a- and empty vector-transfected Cos-7 cells either (1) exposed to lamp light (n=2) (Figure 16A), or (2) not exposed to lamp light (n=1) (Figure 16B). Data are presented as mean \pm SD.

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Figures 17A and 17B

Antagonism by β -ionone (10 μ M) of all-trans-retinal-induced intracellular Ca²+ mobilization in human SNORF36a-transfected Cos-7 cells (n=2) (Figure 17A). Antagonism by β -ionone (10 μ M) of 9-cis-retinal-induced intracellular Ca²+ mobilization in human SNORF36a-transfected Cos-7 cells (n=2) (Figure 17B). The response was measured using Fluorometric Imaging Plate Reader (FLIPRTM). Data are presented as mean \pm SD.

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Figures 18A-18C

Light sensitivity of SNORF36a expressed in oocytes. Oocytes shown in Figure 18A and Figure 18B were injected with 10 ng SNORF36a synthetic mRNA. The current is expressed in nA (nanoampere) while time is expressed in s (second).

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Figure 18A: Current elicited by light exposure (bar) in a voltage-clamped oocyte expressing SNORF36a. The oocyte was pre-incubated with all-trans-retinal (100 nM) in the dark for 24 hours.

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Figure 18B: Response of a second oocyte pre-incubated with all-trans-retinal that had been previously exposed to room light.

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Figure 18C: Control uninjected oocyte, pre-incubated with

PCT/US00/12065

-29-

all-trans-retinal (100 nM) for 24 h, fails to respond to light.

Figure 19

Summary of light sensitivity of oocytes expressing SNORF36 pre-incubated with all-trans-retinal, 13-cis-retinal and ATRA (all-trans retinoic acid). Numbers in parentheses are the numbers of oocytes used.

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DETAILED DESCRIPTION OF THE INVENTION

This invention provides a recombinant nucleic acid comprising a nucleic acid encoding a mammalian SNORF36a receptor, wherein the mammalian receptor-encoding nucleic acid hybridizes under high stringency conditions to a nucleic acid encoding a human SNORF36a receptor and having a sequence identical to the sequence of the human SNORF36a receptor-encoding nucleic acid contained in plasmid pcDNA3.1-hSNORF36a-f (ATCC Patent Depository No. 203977).

This invention further provides a recombinant nucleic acid comprising a nucleic acid encoding a human SNORF36a receptor, wherein the human SNORF36a receptor comprises an amino acid sequence identical to the sequence of the human SNORF36a receptor encoded by the shortest open reading frame indicated in Figures 1A-1C (SEQ ID NO: 1).

This invention provides a recombinant nucleic acid comprising a nucleic acid encoding a mammalian SNORF36b receptor, wherein the mammalian receptor-encoding nucleic acid hybridizes under high stringency conditions to a nucleic acid encoding a human SNORF36b receptor and having a sequence identical to the sequence of the human SNORF36b receptor-encoding nucleic acid contained in plasmid pcDNA3.1-hSNORF36b-f (ATCC Patent Depository No. 203976).

This invention also provides a recombinant nucleic acid comprising a nucleic acid encoding a human SNORF36b receptor, wherein the human SNORF36b receptor comprises an amino acid sequence identical to the sequence of the human SNORF36b receptor encoded by the shortest open reading frame indicated in Figures 3A-3C (SEQ ID NO: 3).

This invention also contemplates recombinant nucleic acids

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which comprise nucleic acids encoding naturally occurring allelic variants of the above. For example, one such allelic variant involves changing Guanine (G) to Adenine (A) at position 39 in Figures 1A-1C (SEQ ID NO: 1). Another example of such an allelic variant involves changing Guanine (G) to Adenine (A) at position 39 in Figures 3A-3C (SEQ ID NO: 3).

The plasmid pcDNA3.1-hSNORF36a-f and plasmid pcDNA3.1-hSNORF36b-f were both deposited on April 28, 1999, with the American Type Culture Collection (ATCC), 10801 University Blvd., Manassas, Virginia 20110-2209, U.S.A. under the provisions of the Budapest Treaty for the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedure and were accorded ATCC Patent Depository Nos. 203977 and 203976, respectively.

The plasmid pEXJ.T3T7-rSNORF36p was deposited on August 17, 1999, with the American Type Culture Collection (ATCC), 10801 University Blvd., Manassas, Virginia 20110-2209, U.S.A. under the provisions of the Budapest Treaty for the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedure and were accorded ATCC Patent Depository No. PTA-534.

The plasmid pEXJ.T7-rSNORF36-f was deposited on January 18, 2000, with the American Type Culture Collection (ATCC), 10801 University Blvd., Manassas, Virginia 20110-2209, U.S.A. under the provisions of the Budapest Treaty for the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedure and were accorded ATCC Patent Depository No. PTA-1216.

Hybridization methods are well known to those of skill in the art. For purposes of this invention, hybridization under high stringency conditions means hybridization performed at 40°C in a hybridization buffer containing 50% formamide, 5X SSC,

-32-

7mM Tris, 1X Denhardt's, 25µg/ml salmon sperm DNA; wash at 50°C in 0.1X SSC, 0.1%SDS.

Throughout this application, the following standard abbreviations are used to indicate specific nucleotide bases:

A = adenine

G = guanine

C = cytosine

T = thymine

R = adenine or guanine

W = adenine or thymine

S = cytosine or guanine

Y = cytosine or thymine

15 K = guanine or thymine

V = adenine, cytosine, or guanine (not thymine)

H = adenine, cytosine, or thymine (not cytosine)

B = cytosine, guanine, or thymine (not adenine)

N = adenine, cytosine, guanine, or thymine (or other modified base such as inosine)

I = inosine

Furthermore, the term "agonist" is used throughout this application to indicate any peptide or non-peptidyl compound which increases the activity of any of the polypeptides of the subject invention. The term "antagonist" is used throughout this application to indicate any peptide or non-peptidyl compound which decreases the activity of any of the polypeptides of the subject invention.

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Furthermore, as used herein, the phrase "pharmaceutically acceptable carrier" means any of the standard pharmaceutically acceptable carriers. Examples include, but are not limited to, phosphate buffered saline, physiological saline, water, and emulsions, such as oil/water emulsions.

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It is possible that the mammalian SNORF36 receptor gene contains introns and furthermore, the possibility exists that additional introns could exist in coding or non-coding regions. In addition, spliced form(s) of mRNA may encode additional amino acids either upstream of the currently defined starting methionine or within the coding region. Further, the existence and use of alternative exons is possible, whereby the mRNA may encode different amino acids within the region comprising the exon. In addition, single amino acid substitutions may arise via the mechanism of RNA editing such that the amino acid sequence of the expressed protein is different than that encoded by the original gene. (Burns, et al., 1996; Chu, et al., 1996). Such variants may exhibit pharmacologic properties differing from the polypeptide encoded by the original gene.

This invention provides splice variants of the mammalian SNORF36 receptors disclosed herein. This invention further provides for alternate translation initiation sites and alternately spliced or edited variants of nucleic acids encoding the SNORF36 receptors of this invention.

This invention also contemplates recombinant nucleic acids which comprise nucleic acids encoding naturally occurring allelic variants of the SNORF36 receptors disclosed herein.

The nucleic acids of the subject invention also include nucleic acid analogs of the human SNORF36a receptor genes, wherein the human SNORF36a receptor gene comprises the nucleic acid sequence shown in Figures 1A-1C (SEQ ID NO: 1) or contained in plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. 203977). Nucleic acid analogs of the human SNORF36a receptor genes differ from the human SNORF36a receptor genes described herein in terms of the identity or location of one or more nucleic acid bases (deletion analogs containing less than all of the nucleic acid bases shown in Figures 1A-1C or

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-34-

contained in plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. 203977), substitution analogs wherein one or more nucleic acid bases shown in Figures 1A-1C or contained in plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. 203977), replaced by other nucleic acid bases, and addition analogs, wherein one or more nucleic acid bases are added to a terminal or medial portion of the nucleic acid sequence) and which encode proteins which share some or all of the properties of the proteins encoded by the nucleic acid sequences shown in Figures 1A-1C or contained in plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. 203977). embodiment of the present invention, the nucleic acid analog encodes a protein which has an amino acid sequence identical to that shown in Figures 2A-2B or encoded by the nucleic acid sequence contained in pcDNA3.1-hSNORF36a-f (ATCC Accession No. 203977). In another embodiment, the nucleic acid analog encodes a protein having an amino acid sequence which differs from the amino acid sequences shown in Figures 2A-2B or encoded by the nucleic acid contained in plasmid pcDNA3.1-No. 203977). hSNORF36a-f (ATCC Accession embodiment, the protein encoded by the nucleic acid analog has a function which is the same as the function of the receptor proteins having the amino acid sequence shown in Figures 2A-2B. In another embodiment, the function of the protein encoded by the nucleic acid analog differs from the function of the receptor protein having the amino acid sequence shown in Figures 2A-2B. In another embodiment, the variation in the nucleic acid sequence occurs within the transmembrane (TM) region of the protein. In a further embodiment, the variation in the nucleic acid sequence occurs outside of the TM region.

The nucleic acids of the subject invention also include nucleic acid analogs of the human SNORF36b receptor genes, wherein the human SNORF36b receptor gene comprises the nucleic acid sequence shown in Figures 3A-3C (SEQ ID NO: 3)

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or contained in plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession Nucleic acid analogs of the human SNORF36b receptor genes differ from the human SNORF36b receptor genes described herein in terms of the identity or location of one or more nucleic acid bases (deletion analogs containing less than all of the nucleic acid bases shown in Figures 3A-3C or contained in plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. 203976), substitution analogs wherein one or more nucleic acid bases shown in Figures 3A-3C or contained in plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. replaced by other nucleic acid bases, and addition analogs, wherein one or more nucleic acid bases are added to a terminal or medial portion of the nucleic acid sequence) and which encode proteins which share some or all of the properties of the proteins encoded by the nucleic acid sequences shown in Figures 3A-3C or contained in plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. 203976). embodiment of the present invention, the nucleic acid analog encodes a protein which has an amino acid sequence identical to that shown in Figures 4A-4C or encoded by the nucleic acid sequence contained in pcDNA3.1-hSNORF36b-f (ATCC Accession No. 203976). In another embodiment, the nucleic acid analog encodes a protein having an amino acid sequence which differs from the amino acid sequences shown in Figures 4A-4C or encoded by the nucleic acid contained in plasmid pcDNA3.1hSNORF36b-f (ATCC Accession No. 203976). In a further embodiment, the protein encoded by the nucleic acid analog has a function which is the same as the function of the receptor proteins having the amino acid sequence shown in Figures 4A-4C. In another embodiment, the function of the protein encoded by the nucleic acid analog differs from the function of the receptor protein having the amino acid sequence shown in Figures 4A-4C. In another embodiment, the variation in the nucleic acid sequence occurs within the transmembrane (TM) region of the protein. In a further embodiment, the variation in the nucleic acid sequence occurs

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-36-

outside of the TM region.

The nucleic acids of the subject invention also include nucleic acid analogs of the rat SNORF36 receptor genes, wherein the rat SNORF36 receptor gene comprises the nucleic acid sequence shown in Figure 5 or contained in plasmid pEXJ.T3T7-rSNORF36p (ATCC Patent Depository No. PTA-534). Nucleic acid analogs of the rat SNORF36 receptor genes differ from the rat SNORF36 receptor genes described herein in terms of the identity or location of one or more nucleic acid bases deletion analogs containing less than all of the nucleic acid bases shown in Figure 5 or contained in plasmid pEXJ.T3T7rSNORF36p (ATCC Patent Depository No. PTA-534), substitution analogs wherein one or more nucleic acid bases shown in Figure 5 or contained in plasmid pEXJ.T3T7-rSNORF36p (ATCC Patent Depository No. PTA-534), are replaced by other nucleic acid bases, and addition analogs, wherein one or more nucleic acid bases are added to a terminal or medial portion of the nucleic acid sequence) and which encode proteins which share some or all of the properties of the proteins encoded by the nucleic acid sequences shown in Figure 5 or contained in plasmid pEXJ.T3T7-rSNORF36p (ATCC Patent Depository No. PTA-In one embodiment of the present invention, the 534). nucleic acid analog encodes a protein which has an amino acid sequence identical to that shown in Figure 6 or encoded by the nucleic acid sequence contained in plasmid pEXJ.T3T7rSNORF36p (ATCC Patent Depository No. PTA-534). In another embodiment, the nucleic acid analog encodes a protein having an amino acid sequence which differs from the amino acid sequences shown in Figure 6 or encoded by the nucleic acid in plasmid pEXJ.T3T7-rSNORF36p (ATCC Patent contained In a further embodiment, the Depository No. PTA-534). protein encoded by the nucleic acid analog has a function which is the same as the function of the receptor proteins having the amino acid sequence shown in Figure 6. In another embodiment, the function of the protein encoded by the

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PCT/US00/12065

-37-

nucleic acid analog differs from the function of the receptor protein having the amino acid sequence shown in Figure 6. In another embodiment, the variation in the nucleic acid sequence occurs within the transmembrane (TM) region of the protein. In a further embodiment, the variation in the nucleic acid sequence occurs outside of the TM region.

The nucleic acids of the subject invention also include nucleic acid analogs of the rat SNORF36 receptor genes, wherein the rat SNORF36 receptor gene comprises the nucleic acid sequence shown in Figures 9A-9C or contained in plasmid pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216). Nucleic acid analogs of the rat SNORF36 receptor genes differ from the rat SNORF36 receptor genes described herein in terms of the identity or location of one or more nucleic acid bases deletion analogs containing less than all of the nucleic acid bases shown in Figures 9A-9C or contained in plasmid pEXJ.T7-(ATCC rSNORF36-f Patent Depository No. PTA-1216), substitution analogs wherein one or more nucleic acid bases shown in Figures 9A-9C or contained in plasmid pEXJ.T7-(ATCC Patent Depository No. PTA-1216), replaced by other nucleic acid bases, and addition analogs, wherein one or more nucleic acid bases are added to a terminal or medial portion of the nucleic acid sequence) and which encode proteins which share some or all of the properties of the proteins encoded by the nucleic acid sequences shown in Figures 9A-9C or contained in plasmid pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216). In one embodiment of the present invention, the nucleic acid analog encodes a protein which has an amino acid sequence identical to that shown in Figures 10A-10C or encoded by the nucleic acid sequence contained in plasmid pEXJ.T7-rSNORF36-f Patent Depository No. PTA-1216). In embodiment, the nucleic acid analog encodes a protein having an amino acid sequence which differs from the amino acid sequences shown in Figures 10A-10C or encoded by the nucleic

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-38-

acid contained in plasmid pEXJ.T7-rsNoRF36-f (ATCC Patent Depository No. PTA-1216). In a further embodiment, the protein encoded by the nucleic acid analog has a function which is the same as the function of the receptor proteins having the amino acid sequence shown in Figures 10A-10C. In another embodiment, the function of the protein encoded by the nucleic acid analog differs from the function of the receptor protein having the amino acid sequence shown in Figures 10A-10C. In another embodiment, the variation in the nucleic acid sequence occurs within the transmembrane (TM) region of the protein. In a further embodiment, the variation in the nucleic acid sequence occurs outside of the TM region.

This invention provides the above-described isolated nucleic acid, wherein the nucleic acid is DNA. In an embodiment, the DNA is cDNA. In another embodiment, the DNA is genomic DNA. In still another embodiment, the nucleic acid is RNA. Methods for production and manipulation of nucleic acid molecules are well known in the art.

This invention further provides nucleic acid which is degenerate with respect to the DNA encoding any of the polypeptides described herein. In an embodiment, the nucleic acid comprises a nucleotide sequence which is degenerate with respect to the nucleotide sequence shown in Figures 1A-1C (SEQ ID NO: 1) or the nucleotide sequence contained in the plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. 203977), that is, a nucleotide sequence which is translated into the same amino acid sequence. In another embodiment, the nucleic acid comprises a nucleotide sequence which is degenerate with respect to the nucleotide sequence shown in Figures 3A-3C (SEQ ID NO: 3) or the nucleotide sequence contained in the plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. 203976), that is, a nucleotide sequence which is translated into the same amino acid sequence.

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PCT/US00/12065

-39-

In another embodiment, the nucleic acid comprises a nucleotide sequence which is degenerate with respect to the nucleotide sequence shown in Figure 5 (SEQ ID NO: 5) or Figures 9A-9C (SEQ ID NO: 7) or the nucleotide sequence contained in the plasmids pEXJ.T3T7-rSNORF36p (ATCC Patent Depository No. PTA-534) or pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216), that is, a nucleotide sequence which is translated into the same amino acid sequence.

- This invention also encompasses DNAs and cDNAs which encode amino acid sequences which differ from those of the polypeptides of this invention, but which should not produce phenotypic changes.
- Alternately, this invention also encompasses DNAs, cDNAs, and RNAs which hybridize to the DNA, cDNA, and RNA of the subject invention. Hybridization methods are well known to those of skill in the art.
- The nucleic acids of the subject invention also include 20 nucleic acid molecules coding for polypeptide analogs, fragments or derivatives of antigenic polypeptides which differ from naturally-occurring forms in terms of the identity or location of one or more amino acid residues 25 (deletion analogs containing less than all of the residues specified for the protein, substitution analogs wherein one or more residues specified are replaced by other residues and addition analogs wherein one or more amino acid residues is added to a terminal or medial portion of the polypeptides) and which share some or all properties of naturally-occurring 30 forms. These molecules include: the incorporation of codons "preferred" for expression by selected non-mammalian hosts; the provision of sites for cleavage by restriction endonuclease enzymes; and the provision of additional initial, terminal or intermediate DNA sequences that 35 facilitate construction of readily expressed vectors.

-40-

creation of polypeptide analogs is well known to those of skill in the art (Spurney, R. F. et al. (1997); Fong, T.M. et al. (1995); Underwood, D.J. et al. (1994); Graziano, M.P. et al. (1996); Guan X.M. et al. (1995)).

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The modified polypeptides of this invention may be transfected into cells either transiently or stably using methods well-known in the art, examples of which are disclosed herein. This invention also provides for binding assays using the modified polypeptides, in which the polypeptide is expressed either transiently or in stable cell lines. This invention further provides a compound identified using a modified polypeptide in a binding assay such as the binding assays described herein.

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The nucleic acids described and claimed herein are useful for the information which they provide concerning the amino acid sequence of the polypeptide and as products for the large scale synthesis of the polypeptides by a variety of recombinant techniques. The nucleic acid molecule is useful for generating new cloning and expression vectors, transformed and transfected prokaryotic and eukaryotic host cells, and new and useful methods for cultured growth of such host cells capable of expression of the polypeptide and related products.

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This invention also provides an isolated nucleic acid encoding species homologs of the SNORF36 receptor encoded by the nucleic acid sequence shown in Figures 1A-1C (SEQ ID NO: 1) or encoded by the plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. 203977). In one embodiment, the nucleic acid encodes a mammalian SNORF36 receptor homolog which has substantially the same amino acid sequence as does the SNORF36 receptor encoded by the plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. 203977). In another embodiment, the nucleic acid encodes a mammalian SNORF36 receptor homolog

PCT/US00/12065

-41-

which has above 75% amino acid identity to the SNORF36 receptor encoded by the plasmid pcDNA3.1-hSNORF36a-f (ATCC No. 203977); preferably above 85% amino acid identity to the SNORF36 receptor encoded by the plasmid No. 203977); most pcDNA3.1-hSNORF36a-f (ATCC Accession preferably above 95% amino acid identity to the SNORF36 receptor encoded by the plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. 203977). In another embodiment, the mammalian SNORF36 receptor homolog has above 70% nucleic acid identity to the SNORF36 receptor gene contained in plasmid pcDNA3.1hSNORF36a-f (ATCC Accession No. 203977); preferably above 80% nucleic acid identity to the SNORF36 receptor gene contained in the plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. 203977); more preferably above 90% nucleic acid identity to the SNORF36 receptor gene contained in the plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. 203977). Examples of methods for isolating and purifying species homologs are described elsewhere (e.g., U.S. Patent No. 5,602,024, WO94/14957, WO97/26853, WO98/15570).

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invention also provides an isolated nucleic acid encoding species homologs of the SNORF36 receptors encoded by the nucleic acid sequence shown in Figures 3A-3C (SEQ ID NO: 3) or encoded by the plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. 203976). In one embodiment, the nucleic acid encodes a mammalian SNORF36 receptor homolog which has substantially the same amino acid sequence as does the SNORF36 receptor encoded by the plasmid pcDNA3.1-hSNORF36b-f No. 203976). (ATCC Accession In another embodiment, the nucleic acid encodes a mammalian SNORF36 receptor homolog which has above 75% amino acid identity to the SNORF36 receptor encoded by the plasmid pcDNA3.1-hSNORF36b-f (ATCC No. 203976); preferably above 85% amino acid Accession identity to the SNORF36 receptor encoded by the plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. 203976); most preferably above 95% amino acid identity to the SNORF36

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-42-

receptor encoded by the plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. 203976). In another embodiment, the mammalian SNORF36 receptor homolog has above 70% nucleic acid identity to the SNORF36 receptor gene contained in plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. 203976); preferably above 80% nucleic acid identity to the SNORF36 receptor gene contained in the plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. 203976); more preferably above 90% nucleic acid identity to the SNORF36 receptor gene contained in the plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. 203976).

This invention also provides an isolated nucleic acid encoding species homologs of the SNORF36 receptors encoded by the nucleic acid sequence shown in Figure 5 (SEQ ID NO: 5) or encoded by the plasmid pEXJ.T3T7-rSNORF36p (ATCC Patent Depository No. PTA-534). In one embodiment, the nucleic acid encodes a mammalian SNORF36 receptor homolog which has substantially the same amino acid sequence as does the SNORF36 receptor encoded by the plasmid pEXJ.T3T7-rSNORF36p (ATCC Patent Depository No. PTA-534). In another embodiment, the nucleic acid encodes a mammalian SNORF36 receptor homolog which has above 75% amino acid identity to the SNORF36 receptor encoded by the plasmid pEXJ.T3T7-rSNORF36p (ATCC Patent Depository No. PTA-534); preferably above 85% amino acid identity to the SNORF36 receptor encoded by the plasmid pEXJ.T3T7-rSNORF36p (ATCC Patent Depository No. PTA-534); most preferably above 95% amino acid identity to the SNORF36 receptor encoded by the plasmid pEXJ.T3T7-rSNORF36p (ATCC Patent Depository No. PTA-534). In another embodiment, the mammalian SNORF36 receptor homolog has above 70% nucleic acid identity to the SNORF36 receptor gene contained in plasmid pEXJ.T3T7-rSNORF36p (ATCC Patent Depository No. PTA-534); preferably above 80% nucleic acid identity to the SNORF36 receptor gene contained in the plasmid pEXJ.T3T7-rSNORF36p (ATCC Patent Depository No. PTA-534); more preferably above 90% nucleic acid identity to the SNORF36 receptor gene

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PCT/US00/12065

-43-

contained in the plasmid pEXJ.T3T7-rSNORF36p (ATCC Patent Depository No. PTA-534).

This invention also provides an isolated nucleic acid encoding species homologs of the SNORF36 receptors encoded by the nucleic acid sequence shown in Figures 9A-9C (SEQ ID NO: 7) or encoded by the plasmid pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216). In one embodiment, the nucleic acid encodes a mammalian SNORF36 receptor homolog which has substantially the same amino acid sequence as does the SNORF36 receptor encoded by the plasmid pEXJ.T7-rSNORF36-(ATCC Patent Depository No. PTA-1216). In another embodiment, the nucleic acid encodes a mammalian SNORF36 receptor homolog which has above 75% amino acid identity to the SNORF36 receptor encoded by the pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216); preferably above 85% amino acid identity to the SNORF36 receptor encoded by the plasmid pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216); most preferably above 95% amino acid identity to the SNORF36 receptor encoded by the plasmid pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216). In another embodiment, the mammalian SNORF36 receptor homolog has above 70% nucleic acid identity to the SNORF36 receptor gene contained in plasmid pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216); preferably above 80% nucleic acid identity to the SNORF36 receptor gene contained in the plasmid pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216); more preferably above 90% nucleic acid identity to the SNORF36 receptor gene contained in the plasmid pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216).

This invention provides an isolated nucleic acid encoding a modified mammalian SNORF36 receptor, which differs from a mammalian SNORF36 receptor by having an amino acid(s) deletion, replacement, or addition in the third intracellular domain.

-44-

This invention provides an isolated nucleic acid encoding a mammalian SNORF36 receptor. In one embodiment, the nucleic acid is DNA. In another embodiment, the DNA is cDNA. In another embodiment, the DNA is genomic DNA. In another embodiment, the nucleic acid is RNA.

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In another embodiment, the mammalian SNORF36 receptor is a human SNORF36a receptor. In another embodiment, the human SNORF36a receptor has an amino acid sequence identical to that encoded by the pcDNA3.1-hSNORF36a-f (ATCC Accession No. 203977). In another embodiment, the human SNORF36a receptor has an amino acid sequence identical to the amino acid sequence shown in Figures 2A-2B (SEQ ID NO: 2).

In another embodiment, the mammalian SNORF36 receptor is a human SNORF36b receptor. In another embodiment, the human SNORF36b receptor has an amino acid sequence identical to that encoded by the plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. 203976). In another embodiment, the human SNORF36b receptor has an amino acid sequence identical to the amino acid sequence shown in Figures 4A-4C (SEQ ID NO: 4).

In an embodiment, the mammalian SNORF36 receptor is a rat SNORF36 receptor. In another embodiment, the rat SNORF36 receptor has an amino acid sequence identical to that encoded by the plasmid pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216). In another embodiment, the rat SNORF36 receptor has an amino acid sequence identical to the amino acid sequence shown in Figures 10A-10C (SEQ ID NO: 8).

This invention provides a purified mammalian SNORF36 receptor protein. In one embodiment, the SNORF36 receptor protein is a human SNORF36a receptor protein. In another embodiment, the SNORF36 receptor protein is a human SNORF36b receptor protein. In a further embodiment, the SNORF36 receptor protein is a rat SNORF36 receptor protein.

-45-

This invention provides a vector comprising the nucleic acid of this invention. This invention further provides a vector adapted for expression in a cell which comprises the regulatory elements necessary for expression of the nucleic acid in the cell operatively linked to the nucleic acid encoding the receptor so as to permit expression thereof, wherein the cell is a bacterial, amphibian, yeast, insect or mammalian cell. In one embodiment, the vector is a baculovirus. In another embodiment, the vector is a plasmid.

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This invention provides a plasmid designated pcDNA3.1-hSNORF36a-f (ATCC Accession No. 203977). This invention also provides a plasmid designated pcDNA3.1-hSNORF36b-f (ATCC Accession No. 203976). This invention provides a plasmid designated pEXJ.T3T7-rSNORF36p (ATCC Patent Depository No. PTA-534). This invention provides a plasmid designated pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216).

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This invention further provides for any vector or plasmid which comprises modified untranslated sequences, which are beneficial for expression in desired host cells or for use in binding or functional assays. For example, a vector or plasmid with untranslated sequences of varying lengths may express differing amounts of the polypeptide depending upon the host cell used. In an embodiment, the vector or plasmid comprises the coding sequence of the polypeptide and the regulatory elements necessary for expression in the host cell.

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This invention provides for a cell comprising the vector of this invention. In one embodiment, the cell is a non-mammalian cell. In one embodiment, the non-mammalian cell is a Xenopus oocyte cell or a Xenopus melanophore cell. In another embodiment, the cell is a mammalian cell. In another embodiment, the cell is a COS-7 cell, a 293 human embryonic

PCT/US00/12065

WO 00/66630

-46-

kidney cell, a NIH-3T3 cell, a LM(tk-) cell, a mouse Y1 cell, or a CHO cell. In another embodiment, the cell is an insect cell. In another embodiment, the insect cell is an Sf9 cell, an Sf21 cell or a Trichoplusia ni 5B-4 cell.

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This invention provides a membrane preparation isolated from the cell in accordance with this invention.

Furthermore, this invention provides for a nucleic acid probe

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comprising at least 15 nucleotides, which probe specifically hybridizes with a nucleic acid encoding a mammalian SNORF36 receptor, wherein the probe has a sequence complementary to a unique sequence present within one of the two strands of the nucleic acid encoding the mammalian SNORF36 receptor contained in plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. 203977), plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. 203976), plasmid pEXJ.T3T7-rSNORF36p (ATCC Patent Depository No. PTA-534) or plasmid pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216).

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This invention further provides a nucleic acid probe comprising at least 15 nucleotides, which probe specifically hybridizes with a nucleic acid encoding a mammalian SNORF36 receptor, wherein the probe has a sequence complementary to a unique sequence present within (a) the nucleic acid sequence shown in Figures 1A-1C (SEQ ID NO: 1) or (b) the reverse complement thereof. This invention further provides a nucleic acid probe comprising at least 15 nucleotides, which probe specifically hybridizes with a nucleic acid encoding a mammalian SNORF36 receptor, wherein the probe has a sequence complementary to a unique sequence present within (a) the nucleic acid sequence shown in Figures 3A-3C (SEQ ID NO: 3) or (b) the reverse complement thereof. This invention also provides a nucleic acid probe comprising at least 15 nucleotides, which probe specifically hybridizes with a nucleic acid encoding a mammalian SNORF36 receptor, wherein

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PCT/US00/12065

-47-

the probe has a sequence complementary to a unique sequence present within (a) the nucleic acid sequence shown in Figure 5 (SEQ ID NO: 5) or (b) the reverse complement thereof. This invention also provides a nucleic acid probe comprising at least 15 nucleotides, which probe specifically hybridizes with a nucleic acid encoding a mammalian SNORF36 receptor, wherein the probe has a sequence complementary to a unique sequence present within (a) the nucleic acid sequence shown in Figures 9A-9C (SEQ ID NO: 7) or (b) the reverse complement thereof. In one embodiment, the nucleic acid is DNA. In another embodiment, the nucleic acid is RNA.

As used herein, the phrase "specifically hybridizing" means the ability of a nucleic acid molecule to recognize a nucleic acid sequence complementary to its own and to form double-helical segments through hydrogen bonding between complementary base pairs.

The nucleic acids of this invention may be used as probes to obtain homologous nucleic acids from other species and to detect the existence of nucleic acids having complementary sequences in samples.

The nucleic acids may also be used to express the receptors they encode in transfected cells.

The use of a constitutively active receptor encoded by SNORF36 either occurring naturally without further modification or after appropriate point mutations, deletions or the like, allows screening for antagonists and *in vivo* use of such antagonists to attribute a role to receptor SNORF36 without prior knowledge of the endogenous ligand.

Use of the nucleic acids further enables elucidation of possible receptor diversity and of the existence of multiple

-48-

subtypes within a family of receptors of which SNORF36 is a member.

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Finally, it is contemplated that this receptor will serve as a valuable tool for designing drugs for treating various pathophysiological conditions such as chronic and acute inflammation, arthritis, autoimmune diseases, transplant rejection, graft vs. host disease, bacterial, protozoan and viral infections, septicemia, AIDS, pain, psychotic and neurological disorders, including anxiety, depression, schizophrenia, dementia, mental retardation, epilepsy, neuromotor disorders, locomotor memory loss, problems, respiratory disorders, asthma, eating/body weight disorders including obesity, bulimia, diabetes, anorexia, and hypotension, vascular hypertension, cardiovascular disorders, ischemia, stroke, cancers, ulcers, urinary retention, sexual/reproductive disorders, circadian rhythm disorders, renal disorders, bone diseases including osteoporosis, benign prostatic hypertrophy, gastrointestinal disorders, nasal congestion, dermatological disorders such as psoriasis, allergies, Parkinson's disease, Alzheimer's disease, acute heart failure, angina disorders, delirium, dyskinesias such as Huntington's disease or Gille's de la Tourette's syndrome, among others and diagnostic assays for This receptor may also serve as a valuable such conditions. tool for designing drugs for chemoprevention.

Methods of transfecting cells e.g. mammalian cells, with such nucleic acid to obtain cells in which the receptor is expressed on the surface of the cell are well known in the for example, U.S. Patent Nos. 5,053,337; 5,516,653; 5,472,866; 5,476,782; 5,360,735; 5,155,218; 5,639,652; 5,602,024; 5,595,880; 5,556,753; 5,545,549; 5,652,113; 5,661,024; 5,766,879; 5,786,155; and 5,786,157, the disclosures of which are hereby incorporated by reference in their entireties into this application.)

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-49-

Such transfected cells may also be used to test compounds and screen compound libraries to obtain compounds which bind to the SNORF36 receptor, as well as compounds which activate or inhibit activation of functional responses in such cells, and therefore are likely to do so in vivo. (See, for example, U.S. Patent Nos. 5,053,337; 5,155,218; 5,360,735; 5,472,866; 5,476,782; 5,516,653; 5,545,549; 5,556,753; 5,595,880; 5,602,024; 5,639,652; 5,652,113; 5,661,024; 5,766,879; 5,786,155; and 5,786,157, the disclosures of which are hereby incorporated by reference in their entireties into this application.)

This invention further provides an antibody capable of binding to a mammalian SNORF36 receptor encoded by a nucleic acid encoding a mammalian receptor. In one embodiment, the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor. In a further embodiment, the mammalian SNORF36 receptor is a rat SNORF36 receptor.

- This invention also provides an agent capable of competitively inhibiting the binding of the antibody to a mammalian SNORF36 receptor. In one embodiment, the antibody is a monoclonal antibody or antisera.
- 25 Methods of preparing and employing antisense oligonucleotides, antibodies, nucleic acid probes transgenic animals directed to the SNORF36 receptor are well known in the art. (See, for example, U.S. Patent Nos. 5, 155, 218; 5,360,735; 5,472,866; 5,053,337; 5,476,782; 5,602,024; 30 5,516,653; 5,545,549; 5,556,753; 5,595,880; 5,639,652; 5,652,113; 5,661,024; 5,766,879; 5,786,155; and 5,786,157, the disclosures of which are hereby incorporated by reference in their entireties into this application.)
- This invention provides for an antisense oligonucleotide having a sequence capable of specifically hybridizing to RNA

PCT/US00/12065

-50-

encoding a mammalian SNORF36 receptor, so as to prevent translation of such RNA. This invention further provides for an antisense oligonucleotide having a sequence capable of specifically hybridizing to genomic DNA encoding a mammalian SNORF36 receptor, so as to prevent transcription of such genomic DNA. In one embodiment, the oligonucleotide comprises chemically modified nucleotides or nucleotide analogues.

This invention also provides for an antibody capable of binding to a mammalian SNORF36 receptor encoded by a nucleic acid in accordance with this invention. In an embodiment, the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor. In a further embodiment, the mammalian SNORF36 receptor is a rat SNORF36 receptor.

Moreover, this invention provides an agent capable of competitively inhibiting the binding of an antibody in accordance with this invention to a mammalian SNORF36 receptor. In one embodiment, the antibody is a monoclonal antibody or antisera.

This invention still further provides a pharmaceutical composition comprising (a) an amount of an oligonucleotide in accordance with this invention capable of passing through a cell membrane and effective to reduce expression of a mammalian SNORF36 receptor and (b) a pharmaceutically acceptable carrier capable of passing through the cell membrane.

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In one embodiment, the oligonucleotide is coupled to a substance which inactivates mRNA. In another embodiment, the substance which inactivates mRNA is a ribozyme. In another embodiment, the pharmaceutically acceptable carrier comprises a structure which binds to a mammalian SNORF36 receptor on a cell capable of being taken up by the cells after binding

PCT/US00/12065

-51-

to the structure. In another embodiment, the pharmaceutically acceptable carrier is capable of binding to a mammalian SNORF36 receptor which is specific for a selected cell type.

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This invention also provides a pharmaceutical composition which comprises an amount of an antibody in accordance with this invention effective to block binding of a ligand to a human SNORF36a receptor or a human SNORF36b receptor and a pharmaceutically acceptable carrier.

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This invention further provides a transgenic, nonhuman mammal expressing DNA encoding a mammalian SNORF36 receptor in accordance with this invention. This invention provides a transgenic, nonhuman mammal comprising а homologous recombination knockout οf а native mammalian receptor. This invention further provides a transgenic, nonhuman mammal whose genome comprises antisense complementary to DNA encoding a mammalian SNORF36 receptor in accordance with this invention so placed within such genome as to be transcribed into antisense mRNA which is complementary and hybridizes with mRNA encoding the mammalian SNORF36 receptor so as to thereby reduce translation or such mRNA and expression of such receptor. In one embodiment, the DNA encoding the mammalian SNORF36 receptor additionally comprises an inducible promoter. In another embodiment, the DNA encoding the mammalian SNORF36 receptor additionally comprises tissue specific regulatory elements. embodiment, the transgenic, nonhuman mammal is a mouse.

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This invention provides for a process for identifying a chemical compound which specifically binds to a mammalian SNORF36 receptor which comprises contacting cells containing DNA encoding, and expressing on their cell surface, the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with the

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PCT/US00/12065

-52-

compound under conditions suitable for binding, and detecting specific binding of the chemical compound to the mammalian This invention further provides for a SNORF36 receptor. for identifying a chemical compound specifically binds to a mammalian SNORF36 receptor which comprises contacting a membrane preparation from cells containing DNA encoding and expressing on their cell surface the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with the compound under conditions suitable for binding, and detecting specific binding of the chemical compound to the mammalian SNORF36 receptor.

In an embodiment, the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor. In another embodiment, the mammalian SNORF36 receptor has substantially the same amino acid sequence as the human SNORF36a receptor encoded by plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. In another embodiment, the mammalian SNORF36 receptor has substantially the same amino acid sequence as the human SNORF36b receptor encoded by plasmid pcDNA3.1hSNORF36b-f (ATCC Accession No. 203976). In another embodiment, the mammalian SNORF36 receptor has substantially the same amino acid sequence as that shown in Figures 2A-2B (SEQ ID NO: 2) or Figures 4A-4C (SEQ ID NO: 4). embodiment, the mammalian SNORF36 receptor has the amino acid sequence shown in Figures 2A-2B (SEQ ID NO: 2) or Figures 4A-4C (SEQ ID NO: 4).

In another embodiment, the mammalian SNORF36 receptor is a rat SNORF36 receptor. In another embodiment, the mammalian SNORF36 receptor has substantially the same amino acid sequence as the rat SNORF36 receptor encoded by plasmid pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216).

In another embodiment, the mammalian SNORF36 receptor has substantially the same amino acid sequence as that shown in

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PCT/US00/12065

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Figures 10A-10C (SEQ ID NO: 8). In another embodiment, the mammalian SNORF36 receptor has the amino acid sequence shown in Figures 10A-10C (SEQ ID NO: 8).

In one embodiment, the compound is not previously known to bind to a mammalian SNORF36 receptor. In one embodiment, the cell is an insect cell. In one embodiment, the cell is a mammalian cell. In another embodiment, the cell nonneuronal in origin. In another embodiment, nonneuronal cell is a COS-7 cell, 293 human embryonic kidney cell, a CHO cell, a NIH-3T3 cell, a mouse Y1 cell, or a In another embodiment, the compound is a LM(tk-) cell. compound not previously known to bind to a mammalian SNORF36 receptor. This invention provides a compound identified by the preceding process according to this invention.

This invention still further provides a process involving competitive binding for identifying a chemical compound which specifically binds to a mammalian SNORF36 receptor which comprises separately contacting cells expressing on their cell surface the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with both the chemical compound and a second chemical compound known to bind to the receptor, and with only the second chemical compound, under conditions suitable for binding of such compounds to the receptor, and detecting specific binding of the chemical compound to the mammalian SNORF36 receptor, a decrease in the binding of the second chemical compound to the mammalian SNORF36 receptor in the presence of the chemical compound being tested indicating that such chemical compound binds to the mammalian SNORF36 receptor.

This invention provides a process involving competitive binding for identifying a chemical compound which specifically binds to a mammalian SNORF36 receptor which

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invention.

-54-

comprises separately contacting a membrane preparation from cells expressing on their cell surface the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with both the chemical compound and a second chemical compound known to bind to the receptor, and with only the second chemical compound, under conditions suitable for binding of such compounds to the receptor, and detecting specific binding of the chemical compound to the mammalian SNORF36 receptor, a decrease in the binding of the second chemical compound to the mammalian SNORF36 receptor in the presence of the chemical compound being tested indicating that such chemical compound binds to the mammalian SNORF36 receptor.

In an embodiment of the present invention, the second chemical compound is a retinoic acid derivative. Examples of retinoic acid derivatives include, but are not limited to, all-trans retinoic acid (ATRA), 9-cis-retinal, 13-cis-retinal, and all-trans-retinal.

In one embodiment, the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor. In another embodiment, the mammalian SNORF36 receptor is a rat SNORF36 receptor. In a further embodiment, the cell is an insect cell. In another embodiment, the cell is a mammalian cell. In another embodiment, the cell is nonneuronal in origin. In another embodiment, the nonneuronal cell is a COS-7 cell, 293 human embryonic kidney cell, a CHO cell, a NIH-3T3 cell, a mouse Y1 cell, or a LM(tk-) cell. In another embodiment, the compound is not previously known to bind to a mammalian SNORF36 receptor. This invention provides for a compound identified by the preceding process according to this

This invention provides for a method of screening a plurality

-55-

of chemical compounds not known to bind to a mammalian SNORF36 receptor to identify a compound which specifically binds to the mammalian SNORF36 receptor, which comprises (a) contacting cells transfected with, and expressing, encoding the mammalian SNORF36 receptor with a compound known to bind specifically to the mammalian SNORF36 receptor; (b) contacting the cells of step (a) with the plurality of compounds not known to bind specifically to the mammalian SNORF36 receptor, under conditions permitting binding of compounds known to bind to the mammalian SNORF36 receptor; determining whether the binding of the compound known to bind to the mammalian SNORF36 receptor is reduced in the presence of the plurality of compounds, relative to the binding of the compound in the absence of the plurality of compounds; and if so (d) separately determining the binding to the mammalian SNORF36 receptor of each compound included in the plurality of compounds, so as to thereby identify any compound included therein which specifically binds to the mammalian SNORF36 receptor.

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This invention provides a method of screening a plurality of chemical compounds not known to bind to a mammalian SNORF36 receptor to identify a compound which specifically binds to mammalian SNORF36 the receptor, which comprises contacting a membrane preparation from cells transfected with, and expressing, DNA encoding the mammalian SNORF36 receptor with the plurality of compounds not known to bind specifically to the mammalian SNORF36 receptor under conditions permitting binding of compounds known to bind to the mammalian SNORF36 receptor; (b) determining whether the binding of a compound known to bind to the mammalian SNORF36 receptor is reduced in the presence of the plurality of compounds, relative to the binding of the compound in the absence of the plurality of compounds; and if so separately determining the binding to the mammalian SNORF36 receptor of each compound included in the plurality of

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-56-

compounds, so as to thereby identify any compound included therein which specifically binds to the mammalian SNORF36 receptor.

In one embodiment, the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor. In a further embodiment, the mammalian SNORF36 receptor is a rat SNORF36 receptor. In another embodiment, the cell is a mammalian cell. In another embodiment, the mammalian cell is non-neuronal in origin. In a further embodiment, the non-neuronal cell is a COS-7 cell, a 293 human embryonic kidney cell, a LM(tk-) cell, a CHO cell, a mouse Yl cell, or an NIH-3T3 cell.

This invention also provides a method of detecting expression of a mammalian SNORF36 receptor by detecting the presence of mRNA coding for the mammalian SNORF36 receptor which comprises obtaining total mRNA from the cell and contacting the mRNA so obtained with a nucleic acid probe according to this invention under hybridizing conditions, detecting the presence of mRNA hybridized to the probe, and thereby detecting the expression of the mammalian SNORF36 receptor by the cell.

This invention further provides for a method of detecting the presence of a mammalian SNORF36 receptor on the surface of a cell which comprises contacting the cell with an antibody according to this invention under conditions permitting binding of the antibody to the receptor, detecting the presence of the antibody bound to the cell, and thereby detecting the presence of the mammalian SNORF36 receptor on the surface of the cell.

This invention still further provides a method of determining the physiological effects of varying levels of activity of mammalian SNORF36 receptors which comprises producing a WO 00/66630 PCT/US00/12065

-57-

transgenic, nonhuman mammal in accordance with this invention whose levels of mammalian SNORF36 receptor activity are varied by use of an inducible promoter which regulates mammalian SNORF36 receptor expression.

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This invention additionally provides a method of determining the physiological effects of varying levels of activity of mammalian SNORF36 receptors which comprises producing a panel of transgenic, nonhuman mammals in accordance with this invention each expressing a different amount of mammalian SNORF36 receptor.

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Moreover, this invention provides method for identifying an antagonist capable of alleviating an abnormality wherein the abnormality is alleviated by decreasing the activity of a mammalian SNORF36 receptor comprising administering compound to a transgenic, nonhuman mammal according to this invention, and determining whether the compound alleviates any physiological and/or behavioral abnormality displayed by the transgenic, nonhuman mammal as a result of overactivity of a mammalian SNORF36 receptor, the alleviation of such an abnormality identifying the compound as an antagonist. an embodiment, the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor. In a further embodiment, the mammalian SNORF36 receptor is a rat SNORF36 receptor. The invention also provides an antagonist identified by the preceding method according to invention. This invention further provides a composition, e.g. a pharmaceutical composition comprising an antagonist this invention and а carrier, according to pharmaceutically acceptable carrier. This invention provides a method of treating an abnormality in a subject wherein the abnormality is alleviated by decreasing the activity of a mammalian SNORF36 receptor which comprises administering to the subject an effective amount of the pharmaceutical composition according to this invention so as to thereby

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PCT/US00/12065

-58-

treat the abnormality.

In addition, this invention provides a method for identifying an agonist capable of alleviating an abnormality in a subject wherein the abnormality is alleviated by increasing the comprising activity of а mammalian SNORF36 receptor administering a compound to a transgenic, nonhuman mammal according to this invention, and determining whether the compound alleviates any physiological and/or behavioral abnormality displayed by the transgenic, nonhuman mammal, the alleviation of such an abnormality identifying the compound as an agonist. In an embodiment, the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor. In a further embodiment, the mammalian SNORF36 receptor is a rat SNORF36 receptor. This invention provides an agonist identified by the preceding method according to this invention. This invention provides a composition, e.g. a pharmaceutical composition comprising an agonist identified by a method according to this invention and a carrier, e.g. a pharmaceutically acceptable carrier.

Moreover, this invention provides a method of treating an abnormality in a subject wherein the abnormality is alleviated by increasing the activity of a mammalian SNORF36 receptor which comprises administering to the subject an effective amount of the pharmaceutical composition of this invention so as to thereby treat the abnormality.

Yet further, this invention provides a method for diagnosing a predisposition to a disorder associated with the activity of a specific mammalian allele which comprises: (a) obtaining DNA of subjects suffering from the disorder; (b) performing a restriction digest of the DNA with a panel of restriction enzymes; (c) electrophoretically separating the resulting DNA fragments on a sizing gel; (d) contacting the resulting gel with a nucleic acid probe capable of

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PCT/US00/12065

-59-

specifically hybridizing with a unique sequence included within the sequence of a nucleic acid molecule encoding a mammalian SNORF36 receptor and labeled with a detectable marker; (e) detecting labeled bands which have hybridized to the DNA encoding a mammalian SNORF36 receptor to create a unique band pattern specific to the DNA of subjects suffering from the disorder; (f) repeating steps (a)-(e) with DNA obtained for diagnosis from subjects not yet suffering from the disorder; and (g) comparing the unique band pattern specific to the DNA of subjects suffering from the disorder from step (e) with the band pattern from step (f) for subjects not yet suffering from the disorder so as to determine whether the patterns are the same or different and thereby diagnose predisposition to the disorder if the patterns are the same.

In one embodiment, the disorder is a disorder associated with the activity of a specific mammalian allele is diagnosed.

This invention also provides a method of preparing a purified mammalian SNORF36 receptor according to this invention which comprises: (a) culturing cells which express the mammalian SNORF36 receptor; (b) recovering the mammalian SNORF36 receptor from the cells; and (c) purifying the mammalian SNORF36 receptor so recovered.

This invention further provides a method of preparing a purified mammalian SNORF36 receptor according to this invention which comprises: (a) inserting a nucleic acid encoding the mammalian SNORF36 receptor into a suitable expression vector; (b) introducing the resulting vector into a suitable host cell; (c) placing the resulting host cell in suitable condition permitting the production of the mammalian SNORF36 receptor; (d) recovering the mammalian SNORF36 receptor so produced; and optionally (e) isolating and/or purifying the mammalian SNORF36 receptor so recovered.

-60-

Furthermore, this invention provides a process for determining whether a chemical compound is a mammalian SNORF36 receptor agonist which comprises contacting cells transfected with and expressing DNA encoding the mammalian SNORF36 receptor with the compound under conditions permitting the activation of the mammalian SNORF36 receptor, and detecting any increase in mammalian SNORF36 receptor activity, so as to thereby determine whether the compound is a mammalian SNORF36 receptor agonist.

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This invention also provides a process for determining whether a chemical compound is a mammalian SNORF36 receptor antagonist which comprises contacting cells transfected with and expressing DNA encoding the mammalian SNORF36 receptor with the compound in the presence of a known mammalian SNORF36 receptor agonist, under conditions permitting the activation of the mammalian SNORF36 receptor, and detecting any decrease in mammalian SNORF36 receptor activity, so as to thereby determine whether the compound is a mammalian SNORF36 receptor antagonist.

In an embodiment, the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor. In another embodiment, the mammalian SNORF36 receptor is a rat SNORF36 receptor.

This invention still further provides a composition, for example a pharmaceutical composition, which comprises an amount of a mammalian SNORF36 receptor agonist determined by a process according to this invention effective to increase activity of a mammalian SNORF36 receptor and a carrier, for example, a pharmaceutically acceptable carrier. In one embodiment, the mammalian SNORF36 receptor agonist is not previously known.

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Also, this invention provides a composition, for example a

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-61-

pharmaceutical composition, which comprises an amount of a mammalian SNORF36 receptor antagonist determined by a process according to this invention effective to reduce activity of a mammalian SNORF36 receptor and a carrier, for example, a pharmaceutically acceptable carrier. In one embodiment, the mammalian SNORF36 receptor antagonist is not previously known.

This invention moreover provides a process for determining whether a chemical compound specifically binds to and activates a mammalian SNORF36 receptor, which comprises contacting cells producing a second messenger response and expressing on their cell surface the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with the chemical compound under conditions suitable for activation of the mammalian SNORF36 receptor, and measuring the second messenger response in the presence and in the absence of the chemical compound, a change, e.g. an increase, in the second messenger response in the presence of the chemical compound indicating that the compound activates the mammalian SNORF36 receptor.

In one embodiment, the second messenger response comprises chloride channel activation and the change in second messenger is an increase in the level of chloride current. In another embodiment, the second messenger response comprises change in intracellular calcium levels and the change in second messenger is an increase in the measure of intracellular calcium. In another embodiment, the second messenger response comprises release of inositol phosphate and the change in second messenger is an increase in the level of inositol phosphate. In another embodiment, the second messenger response comprises release of arachidonic acid and the change in second messenger is an increase in the level of arachidonic acid. In yet another embodiment, the second messenger response comprises GTPyS ligand binding and

WO 00/66630 PCT/US00/12065

-62-

the change in second messenger is an increase in GTPyS ligand binding. In another embodiment, the second messenger response comprises activation of MAP kinase and the change in second messenger response is an increase in MAP kinase activation. In a further embodiment, the second messenger response comprises cAMP accumulation and the change in second messenger response is a reduction in cAMP accumulation.

This invention still further provides a process for determining whether a chemical compound specifically binds to and inhibits activation of a mammalian SNORF36 receptor, which comprises separately contacting cells producing a second messenger response and expressing on their cell surface the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with both the chemical compound and a second chemical compound known to activate the mammalian SNORF36 receptor, and with only the second chemical compound, under conditions suitable for activation of the mammalian SNORF36 receptor, and measuring the second messenger response in the presence of only the second chemical compound and in the presence of both the second chemical compound and the chemical compound, a smaller change, e.g. increase, in the second messenger response in the presence of both the chemical compound and the second chemical compound than in the presence of only the second chemical compound indicating that the chemical compound inhibits activation of the mammalian receptor.

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In an embodiment of the present invention, the second chemical compound is a retinoic acid derivative. Examples of retinoic acid derivatives include, but are not limited to, all-trans retinoic acid (ATRA), 9-cis-retinal, 13-cis-retinal, and all-trans-retinal.

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PCT/US00/12065

-63-

In one embodiment, the second messenger response comprises chloride channel activation and the change in second messenger response is a smaller increase in the level of chloride current in the presence of both the chemical compound and the second chemical compound than in the presence of only the second chemical compound. In another embodiment, the second messenger response comprises change in intracellular calcium levels and the change in second messenger response is a smaller increase in the measure of intracellular calcium in the presence of both the chemical compound and the second chemical compound than in the presence of only the second chemical compound. In another embodiment, the second messenger response comprises release of inositol phosphate and the change in second messenger response is a smaller increase in the level of inositol phosphate in the presence of both the chemical compound and the second chemical compound than in the presence of only the second chemical compound.

In one embodiment, the second messenger response comprises activation of MAP kinase and the change in second messenger response is a smaller increase in the level of MAP kinase activation in the presence of both the chemical compound and the second chemical compound than in the presence of only the second chemical compound. In another embodiment, the second messenger response comprises change in cAMP levels and the change in second messenger response is a smaller change in the level of cAMP in the presence of both the chemical compound and the second chemical compound than in the presence of only the second chemical compound. In another embodiment, the second messenger response comprises release of arachidonic acid and the change in second messenger response is an increase in the level of arachidonic acid levels in the presence of both the chemical compound and the second chemical compound than in the presence of only the second chemical compound. In a further embodiment, the

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-64-

second messenger response comprises GTPyS ligand binding and the change in second messenger is a smaller increase in GTPyS ligand binding in the presence of both the chemical compound and the second chemical compound than in the presence of only the second chemical compound.

In one embodiment, the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor. In a further embodiment, the mammalian SNORF36 receptor is a rat SNORF36 receptor. In another embodiment, the cell is an insect cell. In another embodiment, the cell is a mammalian cell. In another embodiment, the mammalian cell is nonneuronal in origin. In another embodiment, the nonneuronal cell is a COS-7 cell, CHO cell, 293 human embryonic kidney cell, NIH-3T3 cell or LM(tk-) cell. In another embodiment, the compound is not previously known to bind to a mammalian SNORF36 receptor.

Further, this invention provides a compound determined by a process according to this invention and a composition, for example, a pharmaceutical composition, which comprises an amount of a mammalian SNORF36 receptor agonist determined to be such by a process according to this invention effective to increase activity of the mammalian SNORF36 receptor and a carrier, for example, a pharmaceutically acceptable carrier. In one embodiment, the mammalian SNORF36 receptor agonist is not previously known.

This invention also provides a composition, for example, a pharmaceutical composition, which comprises an amount of a mammalian SNORF36 receptor antagonist determined to be such by a process according to this invention, effective to reduce activity of the mammalian SNORF36 receptor and a carrier, for example a pharmaceutically acceptable carrier. In one embodiment, the mammalian SNORF36 receptor antagonist is not

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PCT/US00/12065

-65-

previously known.

This invention yet further provides a method of screening a plurality of chemical compounds not known to activate a mammalian SNORF36 receptor to identify a compound which activates the mammalian SNORF36 receptor which comprises: (a) contacting cells transfected with and expressing the mammalian SNORF36 receptor with the plurality of compounds not known to activate the mammalian SNORF36 receptor, under conditions permitting activation of the mammalian SNORF36 receptor; (b) determining whether the activity of the mammalian SNORF36 receptor is increased in the presence of one or more of the compounds; and if so (c) separately determining whether the activation of the mammalian SNORF36 receptor is increased by any compound included in the plurality of compounds, so as to thereby identify each compound which activates the mammalian SNORF36 receptor. one embodiment, the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor. In a further embodiment, the mammalian SNORF36 receptor is a rat SNORF36 receptor.

This invention provides a method of screening a plurality of chemical compounds not known to inhibit the activation of a mammalian SNORF36 receptor to identify a compound which inhibits the activation of the mammalian SNORF36 receptor, which comprises: (a) contacting cells transfected with and expressing the mammalian SNORF36 receptor with the plurality of compounds in the presence of a known mammalian SNORF36 receptor agonist, under conditions permitting activation of the mammalian SNORF36 receptor; (b) determining whether the extent or amount of activation of the mammalian SNORF36 receptor is reduced in the presence of one or more of the compounds, relative to the extent or amount of activation of the mammalian SNORF36 receptor in the absence of such one or more compounds; and if so (c) separately determining whether

WO 00/66630 PCT/US00/12065

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-66-

each such compound inhibits activation of the mammalian SNORF36 receptor for each compound included in the plurality of compounds, so as to thereby identify any compound included in such plurality of compounds which inhibits the activation of the mammalian SNORF36 receptor.

In one embodiment, the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor. In a further embodiment, the mammalian SNORF36 receptor is a rat SNORF36 receptor. In another embodiment, wherein the cell is a mammalian cell. In another embodiment, the mammalian cell is non-neuronal in origin. In another embodiment, the non-neuronal cell is a COS-7 cell, a 293 human embryonic kidney cell, a LM(tk-) cell or an NIH-3T3 cell.

This invention also provides a composition, for example, a pharmaceutical composition, comprising a compound identified by a method according to this invention in an amount effective to increase mammalian SNORF36 receptor activity and a carrier, for example, a pharmaceutically acceptable carrier.

This invention still further provides a composition, for example, a pharmaceutical composition, comprising a compound identified by a method according to this invention in an amount effective to decrease mammalian SNORF36 receptor activity and a carrier, for example, a pharmaceutically acceptable carrier.

30 Furthermore, this invention provides a method of treating an abnormality in a subject wherein the abnormality is alleviated by increasing the activity of a mammalian SNORF36 receptor which comprises administering to the subject a compound which is a mammalian SNORF36 receptor agonist in an amount effective to treat the abnormality. In one embodiment, the abnormality is a regulation of a steroid

-67-

hormone disorder, an epinephrine release disorder, gastrointestinal disorder, a cardiovascular disorder, an electrolyte balance disorder, hypertension, diabetes, a respiratory disorder, asthma, a reproductive disorder, an immune disorder, an endocrine disorder, a musculoskeletal disorder, a neuroendocrine disorder, cognitive disorder, a memory disorder, somatosensory and neurotransmission disorders, metabolic disorders, a motor coordination disorder, a sensory integration disorder, a motor integration disorder, a dopaminergic function disorder, an appetite disorder, such as anorexia or obesity, a sensory transmission disorder, drug addiction, an olfaction disorder, an autonomic nervous system disorder, pain, neuropsychiatric disorders, affective disorder, migraine, circadian disorders, visual disorders, urinary disorders, blood coaqulationrelated disorders, developmental disorders, or ischemiareperfusion injury-related diseases.

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In a further embodiment of the present invention, the abnormality is a gestational abnormality, a sleep disorder such as insomnia, jet lag or shift-related conditions, disorders associated with melatonin release, and disorders associated with choroid plexus function.

This invention additionally provides a method of treating an abnormality in a subject wherein the abnormality is alleviated by decreasing the activity of a mammalian SNORF36 receptor which comprises administering to the subject a compound which is a mammalian SNORF36 receptor antagonist in an amount effective to treat the abnormality. In one embodiment, the abnormality is a regulation of a steroid hormone disorder, an epinephrine release disorder, a gastrointestinal disorder, a cardiovascular disorder, an electrolyte balance disorder, hypertension, diabetes, a respiratory disorder, asthma, a reproductive function disorder, an immune disorder, an endocrine disorder, a

-68-

musculoskeletal disorder, a neuroendocrine disorder, a cognitive disorder, a memory disorder, somatosensory and neurotransmission disorders, metabolic disorders, a motor coordination disorder, a sensory integration disorder, a motor integration disorder, a dopaminergic function disorder, an appetite disorder, such as anorexia or obesity, a sensory transmission disorder, drug addiction, an olfaction disorder, an autonomic nervous system disorder, pain, neuropsychiatric disorders, affective disorder, migraine, circadian disorders, visual disorders, urinary disorders, blood coagulation-related disorders, developmental disorders, or ischemia-reperfusion injury-related diseases.

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In a further embodiment of the present invention, the abnormality is a gestational abnormality, a sleep disorder such as insomnia, jet lag or shift-related conditions, disorders associated with melatonin release, and disorders associated with choroid plexus function.

This invention also provides a process for making a composition of matter which specifically binds to a mammalian SNORF36 receptor which comprises identifying a chemical compound using a process in accordance with this invention and then synthesizing the chemical compound or a novel structural and functional analog or homolog thereof. In one embodiment, the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor. In another embodiment, the mammalian SNORF36 receptor is a rat SNORF36 receptor.

This invention further provides a process for preparing a composition, for example a pharmaceutical composition which comprises admixing a carrier, for example, a pharmaceutically acceptable carrier, and a pharmaceutically effective amount

of a chemical compound identified by a process in accordance with this invention or a novel structural and functional

-69-

analog or homolog thereof. In one embodiment, the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor. In another embodiment, the mammalian SNORF36 receptor is a rat SNORF36 receptor.

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Thus, once the gene for a targeted receptor subtype is cloned, it is placed into a recipient cell which then expresses the targeted receptor subtype on its surface. cell, which expresses a single population of the targeted human receptor subtype, is then propagated resulting in the establishment of a cell line. This cell line, which constitutes a drug discovery system, is used in two different types of assays: binding assays and functional assays. binding assays, the affinity of a compound for both the receptor subtype that is the target of a particular drug discovery program and other receptor subtypes that could be with side effects measured. associated are measurements enable one to predict the potency of a compound, as well as the degree of selectivity that the compound has for the targeted receptor subtype over other receptor subtypes. The data obtained from binding assays also enable chemists to design compounds toward or away from one or more of the relevant subtypes, as appropriate, for optimal therapeutic efficacy. In functional assays, the nature of the response of the receptor subtype to the compound is determined. Data from the functional assays show whether the compound is acting to inhibit or enhance the activity of the receptor subtype, thus enabling pharmacologists to evaluate compounds rapidly at their ultimate human receptor subtypes targets permitting chemists to rationally design drugs that will be more effective and have fewer or substantially less severe side effects than existing drugs.

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Approaches to designing and synthesizing receptor subtypeselective compounds are well known and include traditional medicinal chemistry and the newer technology of combinatorial

-70-

chemistry, both of which are supported by computer-assisted molecular modeling. With such approaches, chemists and pharmacologists use their knowledge of the structures of the targeted receptor subtype and compounds determined to bind and/or activate or inhibit activation of the receptor subtype to design and synthesize structures that will have activity at these receptor subtypes.

Combinatorial chemistry involves automated synthesis of a variety of novel compounds by assembling them using different combinations of chemical building blocks. The use of combinatorial chemistry greatly accelerates the process of generating compounds. The resulting arrays of compounds are called libraries and are used to screen for compounds ("lead compounds") that demonstrate a sufficient level of activity at receptors of interest. Using combinatorial chemistry it is possible to synthesize "focused" libraries of compounds anticipated to be highly biased toward the receptor target of interest.

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Once lead compounds are identified, whether through the use of combinatorial chemistry or traditional medicinal chemistry or otherwise, a variety of homologs and analogs are prepared to facilitate an understanding of the relationship between chemical structure and biological or functional activity. These studies define structure activity relationships which are then used to design drugs with improved potency, selectivity and pharmacokinetic properties. Combinatorial chemistry is also used to rapidly generate a variety of Traditional medicinal structures for lead optimization. chemistry, which involves the synthesis of compounds one at a time, is also used for further refinement and to generate compounds not accessible by automated techniques. Once such drugs are defined the production is scaled up using standard chemical manufacturing methodologies utilized throughout the pharmaceutical and chemistry industry.

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PCT/US00/12065

-71-

This invention will be better understood from the Experimental Details which follow. However, one skilled in the art will readily appreciate that the specific methods and results discussed are merely illustrative of the invention as described more fully in the claims which follow thereafter.

-72-

EXPERIMENTAL DETAILS

Materials and Methods

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Identification of a fragment of the human SNORF36 receptor A human placental genomic phage library (2.75 million recombinants, Stratagene, LaJolla, CA) was screened using ³²P-labeled oligonucleotide probes, RW76, RW77, RW96, RW97, RW98 and RW99, designed against transmembrane (TM) domains III, V and VI of the human serotonin 5-HT1_D receptor. The overlapping oligonucleotide probes were labeled with DNA Polymerase I (Klenow Fragment) and [α-³²P]dATP/dCTP.

Hybridization of nitrocellulose filter overlays of the plates was performed at low stringency: 40°C in a solution containing 25% formamide, 5x SSC (1X SSC is 0.15M sodium chloride, 0.015M sodium citrate), 1x Denhardt's solution (0.02% polyvinylpyrrolindone, 0.02% Ficoll, 0.02% bovine serum albumin), 7 mM Tris and 25 µg/ml sonicated salmon sperm DNA. The filters were washed at 40 °C in 0.1x SSC containing 0.1% sodium dodecyl sulfate and exposed at -70°C to Kodak XAR film in the presence of intensifying screens.

A positive signal on plate 10 was isolated on a secondary plating. A 2.3 kb fragment, from a HindIII/XbaI digest of DNA isolated from this positive, was identified by Southern blot analysis, subcloned into pUC18 (Gibco BRL, Gaithersburg, MD) and used to transform E.Coli XL1 Blue cells (Stratagene, La Jolla, CA). Plasmid DNA from one transformant, K39, was sequenced using the Sanger dideoxy nucleotide chain termination method (Sanger et al., 1977) on denatured doublestranded plasmid templates, using Sequenase (US Biochemical Corp., Cleveland, OH). Analysis of the sequence of K39 revealed TMs III and IV of a novel GPCR with highest homology to adrenergic receptors.

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PCT/US00/12065

-73-

Isolation of a full-length human SNORF36 receptor

To isolate a full-length SNORF36 receptor, pools of a human hippocampal cDNA library were screened by polymerase chain reaction (PCR) using T604, a K39 specific primer, and either T94, a vector primer, or T603, a K39 specific primer. was performed with Tag DNA Polymerase (Roche Molecular Biochemicals, Indianapolis, IN) with the following protocol: 94°C hold for 5 minutes; 40 cycles of 94°C for 2 minute, 72°C for 4 minutes; 10 minute hold at 72°C; 4°C hold until the samples are run on a gel. High stringency hybridization of isolated colonies from two positive pools with T605, a K39specific oligonucleotide probe, and subsequent PCR testing of positive colonies, resulted in the isolation of a single positive clone named TL252. Analysis of the sequence of TL252 revealed that it contained TMI-TMVII, but was missing the NH2 and COOH termini.

termini of SNORF36, To isolate the COOH Rapid Amplification of cDNA Ends (RACE), was performed using the Clontech Marathon cDNA Amplification kit (Clontech, Palo Alto, CA). Using the supplier's protocol, Marathon adapters were ligated onto ds cDNA prepared from human hippocampal polyA+ RNA. The initial PCR was performed with the supplier's Adapter Primer 1 and A48, a forward primer from TMVI of TL252. 2 µls of this initial PCR reaction was re-amplified using the Adaptor Primer 2 and A49, a forward primer from PCR was performed with Advantage Klentag Polymerase (Clontech, Palo Alto, CA) under the following conditions: 30 seconds at 94° C; 30 cycles of 94° C for 30 seconds, 68° C 4 minutes; and 4° C hold until the products were ready for analysis. A 1.5 kb fragment was isolated from an agarose TAE using the QIAQUICK gel extraction kit Chatsworth, CA) and subcloned into the TA cloning vector (Invitrogen, San Diego, CA). One transformant, AB25, was sequenced using the ABI Big Dye cycle sequencing protocol and

-74-

ABI 377 sequencers (ABI, Foster City, CA). Sequences were analyzed using the Wisconsin Package (GCG, Genetics Computer Group, Madison, WI.). Analysis of the sequence of AB25 revealed the COOH terminus and a stop codon approximately 350 bp downstream from TMVII.

To look for a full-length SNORF36 cDNA, pools of a human hippocampal cDNA library were screened by PCR with TL252specific primers BB788 and BB789 using the Expand Long Template PCR System (Boehringer-Mannheim, Indianapolis, IN). Conditions for PCR amplification were as follows: 94°C hold for 5 minutes; 40 cycles of 94°C for 30 seconds, 68°C for 2 minutes; 68°C hold for 5 minutes; 4°C hold until ready for agarose gel electrophoresis. This screen yielded three positive pools. Subsequent high-stringency hybridization of isolated colonies from two of these pools using a $[Y^{-32}P]$ -ATP-labeled SNORF36-specific probe (BB791) resulted in the identification of 2 positive individual colonies, 260.13.1 and 243.33.3. Sequencing of these clones revealed that they were identical and were full-length at the 3' end. while these clones contained some sequence upstream of TMI, they did not contain the initiating methionine. In addition, these clones both contained an 11 amino acid insert in the first intracellular loop that was not present in TL252.

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To identify the NH₂ terminal sequence, 5'RACE was performed on Marathon-Ready human hippocampal cDNA (Clontech, Palo Alto, CA) according to the Marathon cDNA Amplification Kit protocol. The initial PCR was performed with the supplier's Adapter Primer 1 and BB798 a reverse primer from the second intracellular loop of 260.13.1. One µl of this initial PCR reaction was re-amplified using the Adaptor Primer 2 and BB797, a reverse primer from TMI. PCR was performed with Advantage Klentaq Polymerase (Clontech, Palo Alto, CA) under the following conditions: 30 seconds at 94°C; 5 cycles of 94°C for 30 seconds and 72°C for 3 minutes; 5 cycles of 94°C

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PCT/US00/12065

-75-

for 30 seconds and 70°C for 3 minutes; 23 cycles (initial PCR) or 20 cycles (nested PCR) of 94°C for 30 seconds and 68°C for 3 minutes; 68°C hold for 7 minutes, and 4°C hold until the products were ready for analysis. A 500 bp fragment was isolated from a 1% agarose TAE gel using the QIAQUICK gel extraction kit (QIAGEN, Chatsworth, CA) and sequenced using ABI 377 sequencers and BigDye termination cycle sequencing as described above. Analysis of this sequence revealed the NH₂ terminus including 2 methionines 71 and 52 amino acids upstream from TMI.

A full-length cDNA clone for SNORF36a (without the 11 amino acid insert) was constructed as follows. The 5' end of the cDNA was amplified from hippocampal and pituitary cDNAs using Expand Long Template PCR System (Roche Molecular Biochemicals, Indianapolis, IN) and BB812, a forward primer from the 5'UT also incorporating a BamHI restriction site, and BB813, a reverse primer from the third extracellular loop. Conditions for PCR amplification were as follows: 94°C hold for 5 minutes; 37 cycles of 94°C for 30 seconds; 68°C for 2.5 minutes; 68°C hold for 7 minutes; 4°C hold until ready for agarose gel electrophoresis. 1050 bp bands from 9 independent PCR reactions were cut from a 1% agarose gel, purified using the QIAQUICK gel extraction kit (QIAGEN, Chatsworth, CA), subcloned into the TA cloning vector (Invitrogen, San Diego, CA) and sequenced using the ABI Big Dye cycle sequencing protocol and ABI 377 sequencers (ABI, Foster City, CA). Analysis of these sequences revealed the presence of an allelic variation. Nucleotide 39 (Figures 1A-C and 3A-C) is either an adenine or a guanine. One of these PCR products, F9, matched the consensus sequence with the exception of a single conservative nucleotide change. A 1021 bp BamHI/BglI fragment from F9 was then ligated along with a 1180 bp BglI/EcoRI fragment from the human hippocampal library pool 260.13.1 into a BamHI/EcoRI-cut pcDNA3.1

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PCT/US00/12065

-76-

(Invitrogen, San Diego, CA), and this construct was named The single point mutation introduced by the PCR fragment was corrected with the QuikChange Site-Directed Mutagenesis Kit (Stratagene, LaJolla, CA) using BB926 and The resulting BB927 and the manufacturer's instructions. SNORF36a construct, BO109, was sequenced on both strands as described above. This clone was renamed pcDNA3.1-hSNORF36a-A full-length clone for SNORF36b (with the 11 amino acid insert) was constructed as follows. A 1260 bp fragment was obtained by amplifying BO108 with BB796, a forward primer in TMI and BB936, a reverse primer at the stop codon also incorporating a HindIII site, with the following protocol: 94°C hold for 5 minutes; 32 cycles of 94°C for 30 seconds; 68°C for 2.5 minutes; 68°C hold for 7 minutes; 4°C hold until ready for agarose gel electrophoresis. NspI/HindIII fragment from this PCR product and a 390 bp BamHI/NspI fragment from K81, one of the PCR products from the BB812-BB813 PCR described above, was ligated into a BamHI/HindIII-cut pcDNA3.1 (Invitrogen, San Diego, CA). The resulting SNORF36b construct, BO110, was sequenced on both strands as described above. This clone was renamed pcDNA3.1hsnorF36b-f.

Isolation of a Fragment of the Rat Homologue of SNORF36

To obtain a fragment of the rat homologue of SNORF36, 100 ng of rat genomic DNA (Clontech, Palo Alto, CA) was amplified with BB788, a forward PCR primer corresponding to TMIII of human SNORF36 and BB1097, a reverse primer corresponding to TMV of human SNORF36. PCR was performed with the Expand Long Template PCR System (Roche Molecular Biochemicals) under the following conditions: 30 seconds at 94°C, 45 seconds at 45°C or 50°C, 1.5 minutes at 68°C for 40 cycles, with a pre- and post-incubation of 5 minutes at 94°C and 7 minutes at 68°C respectively. Bands of 800 bp from 7 independent PCR reactions were isolated from a TAE gel, purified using the

-77-

QIAQUICK gel extraction kit (QIAGEN, Chatsworth, CA), and sequenced on both strands as described above. Analysis of the sequence revealed an intron in the second extracellular loop. The consensus of the 7 sequences was used to design forward and reverse PCR primers (BB1182, also incorporating a BamHI restriction site, and BB1183, also incorporating a HindIII site) which were used to amplify a band from rat spinal cord cDNA using the following conditions: 30 seconds at 94°C, 30 seconds at 64°C, 1.5 minutes at 68°C for 37 cycles, with a pre- and post-incubation of 5 minutes at 94°C and 7 minutes at 68°C respectively. Products from 3 independent PCR reactions were digested with BamHI and HindIII and fragments of 250 bp were gel-purified and ligated into the expression vector pEXJ.T3T7. One transformant from each PCR reaction was sequenced as above. The nucleotide sequences of two products were identical to the consensus and one, KO56, was renamed pEXJ-rSNORF36-p.

Isolation of a full-length rat SNORF36 receptor

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To look for the full-length rat SNORF36 cDNA, pools of a rat spinal cord cDNA library were screened by PCR with BB1182, a forward primer from TMIII of the rat SNORF36 fragment, and BB1183, a reverse primer from TMV of the rat SNORF36 fragment. PCR was performed using the Expand Long Template PCR System (Roche Molecular Biochemicals, Indianapolis, IN) under the following conditions: 94°C hold for 5 minutes; 40 cycles of 94°C for 30 seconds, 68°C for 5 minutes; 68°C hold 4°C hold until ready for agarose gel for 7 minutes; electrophoresis. This screen yielded 2 positive pools. Subsequent high-stringency hybridization of isolated colonies from 1 of these pools using a $[\gamma^{-32}P]$ -ATP-labeled rat (BB1298) resulted SNORF36-specific probe identification of a positive individual colony, N323.16.1F, Sequencing of KO109 revealed that it renamed KO109. contained an insert of 2.6 kb, including an open reading

-78-

frame of 1422 nucleotides, 148 nucleotides of 5'UT and approximately 1000 nucleotides of 3'UT. This clone was renamed pEXJ.T7-rSNORF36-f.

Isolation of other species homologs of SNORF36 receptor cDNA A nucleic acid sequence encoding a SNORF36 receptor cDNA from other species may be isolated using standard molecular biology techniques and approaches such as those described below:

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Approach #1: A genomic library (e.g., cosmid, phage, P1, BAC, YAC) generated from the species of interest may be screened with a 32P-labeled oligonucleotide probe corresponding to a fragment of the human or rat SNORF36 receptors whose sequence is shown in Figures 1A-1C, 3A-3C, 5 or 9A-9C to isolate a genomic clone. The full-length sequence may be obtained by sequencing this genomic clone. If one or more introns are present in the gene, the full-length intronless gene may be obtained from cDNA using standard molecular For example, a forward PCR primer designed in the 5'UT and a reverse PCR primer designed in the 3'UT may be used to amplify a full-length, intronless receptor from Standard molecular biology techniques could be used to subclone this gene into a mammalian expression vector.

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Approach #2: Standard molecular biology techniques may be used to screen commercial cDNA phage libraries of the species of interest by hybridization under reduced stringency with a ³²P-labeled oligonucleotide probe corresponding to a fragment of the sequences shown in Figures 1A-1C, 3A-3C, 5 or 9A-9C. One may isolate a full-length SNORF36 receptor by obtaining a plaque purified clone from the lambda libraries and then subjecting the clone to direct DNA sequencing. Alternatively, standard molecular biology techniques could be used to screen cDNA plasmid libraries by PCR amplification

-79-

of library pools using primers designed against a partial species homolog sequence. A full-length clone may be isolated by Southern hybridization of colony lifts of positive pools with a ³²P-oligonucleotide probe.

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Approach #3: 3' and 5' RACE may be utilized to generate PCR products from cDNA derived from the species of interest expressing SNORF36 which contain the additional sequence of SNORF36. These RACE PCR products may then be sequenced to determine the additional sequence. This new sequence is then used to design a forward PCR primer in the 5'UT and a reverse primer in the 3'UT. These primers are then used to amplify a full-length SNORF36 clone from cDNA.

Examples of other species include, but are not limited to, mouse, dog, monkey, hamster and guinea pig.

Primers and probes used in the identification of SNORF36:

RW76: 5' - CATCGCCCTCGACGTGCTGCTGCACCTCATCCT TGCACCT -3' (SEQ ID NO: 9)

RW77: 5'- CATGGACAGGTCGCGCTACCGCGTGTCCACGTTCTACC
TACTCCA -3' (SEQ ID NO: 10)

RW96: 5'- GGCATCATCATGGGCACCTTCATCCTCTGCTGGCTGCC
CTTCTTC -3' (SEQ ID NO: 11)

25 RW97: 5'- GCAGAAGGGCAGAACAAGAGCCACGATGAAGAAGGGCA GCCAGCA -3' (SEQ ID NO: 12)

RW98: 5'- TGGCTGTCATCGGACATCACTTGTTGCACTGCCTCCAT

CCTGCAC -3' (SEQ ID NO: 13)

RW99: 5'- GTAGCGGTCCAGGGCGATGACACAGAGGTGCAGGATGG
AGGCAGT -3' (SEQ ID NO: 14)

T604: 5'- CCAGCCGAAGAAGGGTGGCAGACTCCA -3' (SEQ ID NO: 15)

T94: 5'- CTTCTAGGCCTGTACGGAAGTGTTA -3' (SEQ ID NO: 16)

T603: 5'- GCACAGGCTGCGAGTTCTATTCCTT -3' (SEQ ID NO: 17)

35 T605: 5' - CTGGTAATCACACCCGCTGGCCACCTTTGGTGTGGC
GTCCAAG -3' (SEQ ID NO: 18)

-80-

A48: 5'- AGATCATGCTGGTCATCCTCC -3' (SEQ ID NO: 19)

A49: 5'- TCGTGCTCTCCTGGGCTCCCT -3' (SEQ ID NO: 20)

BB788: 5'- TCCTCCATGATCACCCTGACGC -3' (SEQ ID NO: 21)

BB789: 5'- TCTGGAGAGCCCGTCCTGTCTCC -3' (SEQ ID NO: 22)

BB791: 5'- CGGCCGTGCGTGCCTACACCATGCTTCTCTGCTGCTTC
GTGTTCTTCC -3' (SEQ ID NO: 23)

BB798: 5'- TTGGACGCCACACCAAAGGTGGCC -3' (SEQ ID NO: 24)

BB797: 5'- GGTATAGATGACCGTCAGGTTGCC -3' (SEQ ID NO: 25)

BB812: 5'-CGAACAGGATCCTCTCTGTGGGCTCGAGCAAGGACC -3'
(SEQ ID NO: 26)

BB813: 5'- ACGTGTGCGTACCCAGCAAAGGCC -3' (SEQ ID NO: 27)

BB926: 5'- GTCCCACAGCACCTGGGACTTGGGCTGC -3' (SEQ ID NO: 28)

BB927: 5'- GCAGCCCAAGTCCCAGGTGCTGTGGGAC -3' (SEQ ID NO: 29)

BB796: 5'- GGCAACCTGACGGTCATCTATACC -3' (SEQ ID NO: 30)

BB936: 5'- CAGCATAAGCTTCCAGTGGGCGTCCTACATCCTGG -3' (SEQ ID NO: 31)

BB1097: 5'- CAGTAGATGATGATAAGCAGAGG -3' (SEQ ID NO: 32)

BB1182: 5'- CGAACAGGATCCCATAGCCATGGACCGCTATCTGG -3'
(SEQ ID NO: 33)

BB1183: 5'- CCTAGCAAGCTTGAGGAAGAAGACAAAGCAGAAGAGC -3'
(SEO ID NO: 34)

BB1298: 5'-CGGACGCACTAGTCCTGCTAGGTGTCTGGCTCTATGCCCTGG

CCTGG -3' (SEQ ID NO: 35)

Host cells

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A broad variety of host cells can be used to study heterologously expressed proteins. These cells include, but are not limited to, mammalian cell lines such as: COS-7, CHO, LM(tk-), HEK293, etc.; insect cell lines such as Sf9, Sf21, Trichoplusia ni 5B-4, etc.; amphibian cells such as Xenopus oocytes; assorted yeast strains; assorted bacterial cell strains; and others. Culture conditions for each of these cell types are specific and are known to those familiar with

PCT/US00/12065

-81-

the art. The cells used to express human SNORF36 receptor were COS-7 and Human embryonic kidney (HEK) 293 cells.

COS-7 cells are grown on 150 mm plates in DMEM with supplements (Dulbecco's Modified Eagle Medium with 10% bovine calf serum, 4 mM glutamine, 100 units/ml penicillin/100 μ g/ml streptomycin) at 37°C, 5% CO₂. Stock plates of COS-7 cells are trypsinized and split 1:6 every 3-4 days.

HEK293 cells are grown on 150 mm plates in DMEM with supplements (10% bovine calf serum, 4 mM glutamine, 100 units/ml penicillin/100 μ g/ml streptomycin) at 37°C, 5% CO₂. Stock plates of 293 cells are trypsinized and split 1:6 every 3-4 days.

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Transient expression

DNA encoding proteins to be studied can be transiently expressed in a variety of mammalian, insect, amphibian and other cell lines by several methods, such as, calcium phosphate-mediated, DEAE-dextran mediated, liposomal-mediated, viral-mediated, electroporation-mediated and microinjection delivery. Each of these methods may require optimization of assorted experimental parameters depending on the DNA, cell line, and the type of assay to be subsequently employed.

A typical protocol for the DEAE-dextran method as applied to Cos-7 and HEK293 cells is described as follows. Cells to be used for transfection are split 24 hours prior to the transfection to provide flasks which are 70-80% confluent at the time of transfection. Briefly, 8 μ g of receptor DNA plus 8 μ g of any additional DNA needed (e.g. G_{α} protein expression vector, reporter construct, antibiotic resistance marker, mock vector, etc.) are added to 9 ml of complete DMEM plus DEAE-dextran mixture (10 mg/ml in PBS). Cells plated into

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-82-

a T225 flask (sub-confluent) are washed once with PBS and the DNA mixture is added to each flask. The cells are allowed to incubate for 30 minutes at 37°C, 5% CO₂. Following the incubation, 36 ml of complete DMEM with 80 µM chloroquine is added to each flask and allowed to incubate an additional 3 hours. The medium is then aspirated and 24 ml of complete medium containing 10% DMSO for exactly 2 minutes and then aspirated. The cells are then washed 2 times with PBS and 30 ml of complete DMEM added to each flask. The cells are then allowed to incubate over night. The next day the cells are harvested by trypsinization and reseeded into 96 well plates.

Stable expression

Heterologous DNA can be stably incorporated into host cells, causing the cell to perpetually express a foreign protein. Methods for the delivery of the DNA into the cell are similar to those described above for transient expression but require the co-transfection of an ancillary gene to confer drug resistance on the targeted host cell. The ensuing drug resistance can be exploited to select and maintain cells that have taken up the DNA. An assortment of resistance genes are restricted to available including but not purposes of and hygromycin. For kanamycin, concerning the receptor of this invention, stable expression of a heterologous receptor protein is typically carrier out in, mammalian cells including but not necessarily restricted to, CHO, HEK293, LM(tk-), etc. In addition native cell lines that naturally carry and express the nucleic acid sequences for the receptor may be used without the need to engineer the receptor complement.

Functional assays

35 Cells expressing the receptor DNA of this invention may be used to screen for ligands to said receptor using functional

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PCT/US00/12065

-83-

assays. Once a ligand is identified the same assays may be used to identify agonists or antagonists of the receptor that may be employed for a variety of therapeutic purposes. It is well known to those in the art that the over-expression of a GPCR can result in the constitutive activation of intracellular signaling pathways. In the same manner, over-expression of the SNORF36a receptor in any cell line as described above, can result in the activation of the functional responses described below, and any of the assays herein described can be used to screen for agonist, partial agonist, inverse agonist and antagonist ligands of the SNORF36 receptor.

A wide spectrum of assays can be employed to screen for the presence of SNORF36 receptor ligands. These assays range from traditional measurements of total inositol phosphate intracellular levels, cAMP accumulation. mobilization, and potassium currents, for example; to systems measuring these same second messengers but which have been modified or adapted to be of higher throughput, more generic and more sensitive; to cell based assays reporting more general cellular events resulting from receptor activation differentiation, changes, metabolic division/proliferation. Description of several such assays follow.

Cyclic AMP (cAMP) assay

The receptor-mediated stimulation or inhibition of cyclic AMP (cAMP) formation may be assayed in cells expressing the receptors. Cells are plated in 96-well plates or other vessels and preincubated in a buffer such as HEPES buffered saline (NaCl (150 mM), CaCl $_2$ (1 mM), KCl (5 mM), glucose (10 mM)) supplemented with a phosphodiesterase inhibitor such as 5mM theophylline, with or without protease inhibitor cocktail (For example, a typical inhibitor cocktail contains 2 μ g/ml aprotinin, 0.5 mg/ml leupeptin, and 10 μ g/ml phosphoramidon.)

-84-

for 20 min at 37°C, in 5% CO₂. Test compounds are added with or without 10mM forskolin and incubated for an additional 10 min at 37°C. The medium is then aspirated and the reaction stopped by the addition of 100 mM HCl or other methods. The plates are stored at 4°C for 15 min, and the cAMP content in the stopping solution is measured by radioimmunoassay. Radioactivity may be quantified using a gamma counter equipped with data reduction software. Specific modifications may be performed to optimize the assay for the receptor or to alter the detection method of cAMP.

Arachidonic acid release assay

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Cells expressing the receptor are seeded into 96 well plates or other vessels and grown for 3 days in medium with $^{3}\text{H-arachidonic acid (specific activity = 0.75}$ supplements. μCi/ml) is delivered as a 100 μL aliquot to each well and samples are incubated at 37° C, 5% CO₂ for 18 hours. The labeled cells are washed three times with medium. The wells are then filled with medium and the assay is initiated with the addition of test compounds or buffer in a total volume of 250 μ L. Cells are incubated for 30 min at 37°C, 5% CO₂. Supernatants are transferred to a microtiter plate and evaporated to dryness at 75°C in a vacuum oven. Samples are then dissolved and resuspended in 25 μL distilled water. Scintillant (300 µL) is added to each well and samples are counted for 3H in a Trilux plate reader. Data are analyzed using nonlinear regression and statistical techniques available in the GraphPAD Prism package (San Diego, CA).

Intracellular calcium mobilization assays

The intracellular free calcium concentration may be measured by microspectrofluorimetry using the fluorescent indicator dye Fura-2/AM (Bush et al, 1991). Cells expressing the receptor are seeded onto a 35 mm culture dish containing a glass coverslip insert and allowed to achieve overnight. Cells are then washed with HBS and loaded with 100 μ L of

-85-

Fura-2/AM (10 μ M) for 20 to 40 min. After washing with HBS to remove the Fura-2/AM solution, cells are equilibrated in HBS for 10 to 20 min. Cells are then visualized under the 40% objective of a Leitz Fluovert FS microscope and fluorescence emission is determined at 510 nM with excitation wavelengths alternating between 340 nM and 380 nM. Raw fluorescence data are converted to calcium concentrations using standard calcium concentration curves and software analysis techniques.

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In another method, the measurement of intracellular calcium can also be performed on a 96-well (or higher) format and with alternative calcium-sensitive indicators, preferred examples of these are: aequorin, Fluo-3, Fluo-4, Fluo-5, Calcium Green-1, Oregon Green, and 488 BAPTA. After activation of the receptors with agonist ligands the emission elicited by the change of intracellular calcium concentration can be measured by a luminometer, or a fluorescence imager; a preferred example of this is the fluorescence imager plate reader (FLIPR).

Cells expressing the receptor of interest are plated into clear, flat-bottom, black-wall 96-well plates (Costar) at a density of 30,000-80,000 cells per well and allowed to incubate over night at 5% CO₂, 37°C. The growth medium is aspirated and 100 µl of dye loading medium is added to each The loading medium contains: Hank's BSS (without phenol red) (Gibco), 20 mM HEPES (Sigma), 0.1% BSA (Sigma), dye/pluronic acid mixture (e.g. 1 mM Flou-3, AM (Molecular Probes), 10% pluronic acid (Molecular Probes); immediately before use), and 2.5 mΜ probenecid (Sigma) (prepared fresh)). The cells are allowed to incubate for about 1 hour at 5% CO₂, 37°C.

During the dye loading incubation the compound plate is prepared. The compounds are diluted in wash buffer (Hank's

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-86-

BSS without phenol red), 20 mM HEPES, 2.5 mM probenecid to a 3X final concentration and aliquoted into a clear v-bottom plate (Nunc). Following the incubation the cells are washed to remove the excess dye. A Denley plate washer is used to gently wash the cells 4 times and leave a 100 μl final volume of wash buffer in each well. The cell plate is placed in the center tray and the compound plate is placed in the right The FLIPR software is setup for the tray of the FLIPR. experiment, the experiment is run and the data are collected. The data are then analyzed using an excel spreadsheet program.

Antagonist ligands are identified by the inhibition of the signal elicited by agonist ligands.

The intracellular free calcium (Ca^{2+}) concentration may be measured by the Fluorescence Imager Plate Reader (FLIPR $^{\text{TM}}$).

Cells transfected with appropriate DNA as described earlier were plated into clear, flat-bottom, black-wall 96-well plates (Costar) at a density of 80,000-150,000 cells per well and allowed to incubate for 24 hr at 5% CO₂, 37°C. necessary, the cells were placed in a waterbath maintained at 37°C and were exposed to a lamp light (50W) placed at a distance of approximately 30 cm, 90 to 120 min before the 25 imaging. The growth medium was aspirated and 100 μl of loading medium containing fluo-3 dye was added to each well The loading medium contained: 60 min before the imaging. Hank's BSS (without phenol red) (Gibco), 20 mM HEPES (Sigma), 0.1 or 1% BSA (Sigma), dye/pluronic acid mixture (e.g. 1 mM 30 and 10% pluronic acid (Molecular Probes) Flou-3. AΜ (Molecular Probes) mixed immediately before use), and 2.5 mM probenecid (Sigma) (prepared fresh). The cells were allowed to incubate for about 1 hour at 5% CO2, 37°C.

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-87-

medium, the test compound plate was prepared. Since retinoids are chemically unstable and can undergo rapid photoisomerization (Hu et al., 1994), care was taken to weigh, dissolve and prepare appropriate concentrations of the test compounds under photographic darkroom lights. compounds were diluted in wash buffer (Hank's BSS (without phenol red), 20 mM HEPES, 2.5 mM probenecid) to a 4X final concentration and aliquoted into a clear v-bottom plate The test compounds were protected from light till their addition to cells. Following the incubation with the dye-loading medium, the cells were washed to remove the excess dye. A Denley plate washer was used to gently wash the cells 4 times and leave a 100 µl final volume of wash buffer in each well. The cell plate was placed in the center tray and the test compound plate was placed in the right tray of the FLIPR. The FLIPR software was setup for the experiment, the experiment was run and the data were collected. For the agonist experiment, the sampling rate in the FLIPR was every 1 sec for the first minute and every 2 sec for the next two minutes. In this paradigm, the test compound was added after recording base line for the first 10 sec. For the antagonist experiment, the sampling rate in the FLIPR was every 1 sec for the first minute, every 6 sec for the next 5 minutes, every 1 sec for the next one minute followed by every 2 sec next two minutes. To evaluate the antagonistic activity, the test compound was added to the cells 10 sec after commencing the recording and the agonist was added at 310 sec of the recording. Baseline subtraction and negative control corrections were performed on the traces. The collected data were then analyzed using an Excel spreadsheet program.

Inositol phosphate assay

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Human SNORF36a receptor-mediated activation of the inositol phosphate (IP) second messenger pathways was assessed by radiometric measurement of IP products.

-88-

For example, in a 96 well microplate format assay, cells are plated at a density of 70,000 cells per well and allowed to incubate for 24 hours. The cells are then labeled with 0.5 μCi [3H]myo-inositol overnight at 37°C, 5% CO₂. Immediately before the assay, the medium is removed and replaced with 90 uL of PBS containing 10 mM LiCl. The plates are then incubated for 15 min at 37°C, 5% CO₂. Following incubation, the cells are challenged with agonist (10 ul/well; 10x concentration) for 30 min at 37°C, challenge is terminated by the addition of 100 µL of 50% v/v trichloroacetic acid, followed by incubation at 4°C for greater than 30 minutes. Total IPs are isolated from the lysate by ion exchange chromatography. Briefly, the lysed contents of the wells are transferred to a Multiscreen HV filter plate (Millipore) containing Dowex AG1-X8 (200-400 mesh, formate form). The filter plates are prepared adding 100 µL of Dowex AG1-X8 suspension (50% v/v, water: resin) to each well. The filter plates are placed on a vacuum manifold to wash or elute the resin bed. Each well is first washed 2 times with 200 µl of 5 mM myo-inositol. Total [3H]inositol phosphates are eluted with 75 µl of 1.2 M ammonium formate/0.1 M formic acid solution into 96-well plates. uL of scintillation cocktail is added to each well, and the radioactivity is determined by liquid scintillation counting.

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Cells were plated at a density of 70,000 cells per well and allowed to incubate for 24 hours. The cells were then labeled with 0.5 μ Ci [³H]myo-inositol overnight at 37°C, 5% CO₂. Immediately before the assay, the medium was removed and replaced with 180 μ L of Phosphate-Buffered Saline (PBS) containing 10 mM LiCl. The plates were then incubated for 20 min at 37°C, 5% CO₂. Following the incubation, the cells were challenged with agonist (20 μ l/well; 10x concentration) for 30 min at 37°C and were simultaneously either exposed or not to a lamp light (50W) placed at approximately 30 cm distance from the cells. The challenge was terminated by the addition

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PCT/US00/12065

-89-

of 100 μ L of 5% v/v trichloroacetic acid, followed by incubation at 4°C for greater than 30 minutes. Total IPs were isolated from the lysate by ion exchange chromatography. Briefly, the lysed contents of the wells were transferred to a Multiscreen HV filter plate (Millipore) containing Dowex AG1-X8 (200-400 mesh, formate form). The filter plates were prepared adding 100 μ L of Dowex AG1-X8 suspension (50% v/v, water: resin) to each well. The filter plates were placed on a vacuum manifold to wash or elute the resin bed. Each well was first washed 2 times with 200 μ l of 5 mM myoinositol. Total [³H]inositol phosphates were eluted with 75 μ l of 1.2M ammonium formate/0.1M formic acid solution into 96-well plates. 200 μ L of scintillation cocktail was added to each well, and the radioactivity was determined by liquid scintillation counting.

GTPvS functional assay

Membranes from cells expressing the receptor are suspended in assay buffer (e.g., 50 mM Tris, 100 mM NaCl, 5 mM MgCl2, 10 µM GDP, pH 7.4) with or without protease inhibitors (e.g., 0.1% bacitracin). Membranes are incubated on ice for 20 minutes, transferred to a 96-well Millipore microtiter GF/C filter plate and mixed with GTPY35S (e.g., cpm/sample, specific activity ~1000 Ci/mmol) plus or minus unlabeled GTPyS (final concentration = 100 μM). membrane protein concentration ≈ 90 μg/ml. Samples are incubated in the presence or absence of test compounds for 30 min. at room temperature, then filtered on a Millipore vacuum manifold and washed three times with cold (4°C) assay buffer. Samples collected in the filter plate are treated with scintillant and counted for 35S in a Trilux (Wallac) liquid scintillation counter. It is expected that optimal results are obtained when the receptor membrane preparation is derived from an appropriately engineered heterologous expression system, i.e., an expression system resulting in high levels of expression of the receptor and/or expressing

-90-

G-proteins having high turnover rates (for the exchange of GDP for GTP). GTPyS assays are well-known to those skilled in the art, and it is contemplated that variations on the method described above, such as are described by Tian et al. (1994) or Lazareno and Birdsall (1993), may be used.

Microphysiometric assay

Because cellular metabolism is intricately involved in a broad range of cellular events (including receptor activation of multiple messenger pathways), the use of microphysiometric measurements of cell metabolism can in principle provide a generic assay of cellular activity arising from the activation of any orphan receptor regardless of the specifics of the receptor's signaling pathway.

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General guidelines for transient receptor expression, cell preparation and microphysiometric recording are described elsewhere (Salon, J.A. and Owicki, J.A., 1996). Typically cells expressing receptors are harvested and seeded at 3 \times 10⁵ cells per microphysiometer capsule in complete media 24 hours prior to an experiment. The media is replaced with serum free media 16 hours prior to recording to minimize nonspecific metabolic stimulation by assorted and ill-defined On the day of the experiment the cell serum factors. capsules are transferred to the microphysiometer and allowed to equilibrate in recording media (low buffer RPMI 1640, no (Molecular Devices Corporation, bicarbonate, no serum Sunnyvale, CA) containing 0.1% fatty acid free BSA), during which a baseline measurement of basal metabolic activity is established.

A standard recording protocol specifies a 100 μ l/min flow rate, with a 2 min total pump cycle which includes a 30 sec flow interruption during which the acidification rate measurement is taken. Ligand challenges involve a 1 min 20 sec exposure to the sample just prior to the first post

PCT/US00/12065

-91-

challenge rate measurement being taken, followed by two additional pump cycles for a total of 5 min 20 sec sample exposure. Typically, drugs in a primary screen are presented to the cells at 10 µM final concentration.

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Follow up experiments to examine dose-dependency of active compounds are then done by sequentially challenging the cells with a drug concentration range that exceeds the amount needed to generate responses ranging from threshold to maximal levels. Ligand samples are then washed out and the acidification rates reported are expressed as a percentage increase of the peak response over the baseline rate observed just prior to challenge.

15 MAP kinase assay

MAP kinase (mitogen activated kinase) may be monitored to evaluate receptor activation. MAP kinase is activated by multiple pathways in the cell. A primary mode of activation involves the ras/raf/MEK/MAP kinase pathway. Growth factor (tyrosine kinase) receptors feed into this pathway via SHC/Grb-2/SOS/ras. Gi coupled receptors are also known to activate ras and subsequently produce an activation of MAP kinase. Receptors that activate phospholipase C (such as Gg/G11-coupled) produce diacylglycerol (DAG) consequence of phosphatidyl inositol hydrolysis. DAG activates protein kinase C which in turn phosphorylates MAP kinase.

MAP kinase activation can be detected by several approaches. One approach is based on an evaluation of the phosphorylation state, either unphosphorylated (inactive) or phosphorylated (active). The phosphorylated protein has a slower mobility in SDS-PAGE and can therefore be compared with the unstimulated protein using Western blotting. Alternatively, antibodies specific for the phosphorylated protein are available (New England Biolabs) which can be used to detect

-92-

an increase in the phosphorylated kinase. In either method, cells are stimulated with the test compound and then extracted with Laemmli buffer. The soluble fraction is applied to an SDS-PAGE gel and proteins are transferred electrophoretically to nitrocellulose or Immobilon. Immunoreactive bands are detected by standard Western blotting technique. Visible or chemiluminescent signals are recorded on film and may be quantified by densitometry.

Another approach is based on evaluation of the MAP kinase activity via a phosphorylation assay. Cells are stimulated with the test compound and a soluble extract is prepared. The extract is incubated at 30° C for 10 min with gamma- 32 P-ATP, an ATP regenerating system, and a specific substrate for MAP kinase such as phosphorylated heat and acid stable protein regulated by insulin, or PHAS-I. The reaction is terminated by the addition of H₃PO₄ and samples are transferred to ice. An aliquot is spotted onto Whatman P81 chromatography paper, which retains the phosphorylated protein. The chromatography paper is washed and counted for ³²P in a liquid scintillation counter. Alternatively, the cell extract is incubated with gamma-32P-ATP, regenerating system, and biotinylated myelin basic protein bound by streptavidin to a filter support. The myelin basic protein is a substrate for activated MAP kinase. phosphorylation reaction is carried out for 10 min at 30°C. The extract can then by aspirated through the filter, which retains the phosphorylated myelin basic protein. The filter is washed and counted for 32P by liquid scintillation counting.

Cell proliferation assay

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Receptor activation of the orphan receptor may lead to a mitogenic or proliferative response which can be monitored via 3H -thymidine uptake. When cultured cells are incubated with 3H -thymidine, the thymidine translocates into the nuclei

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-93-

where it is phosphorylated to thymidine triphosphate. nucleotide triphosphate is then incorporated into the cellular DNA at a rate that is proportional to the rate of cell growth. Typically, cells are grown in culture for 1-3 days. Cells are forced into quiescence by the removal of serum for 24 hrs. A mitogenic agent is then added to the 24 hrs later, the cells are incubated with ³Hthymidine at specific activities ranging from 1 to 10 µCi/ml Harvesting procedures may hrs. trypsinization and trapping of cells by filtration over GF/C filters with or without a prior incubation in TCA to extract soluble thymidine. The filters are processed with scintillant and counted for ³H by liquid scintillation counting. Alternatively, adherent cells are fixed in MeOH in water, and solubilized TCA, washed deoxycholate/0.1 N NaOH. The soluble extract is transferred scintillation vials and counted for 3H by liquid scintillation counting.

Alternatively, cell proliferation can be assayed by measuring the expression of an endogenous or heterologous gene product, expressed by the cell line used to transfect the orphan receptor, which can be detected by methods such as, but not limited to, florescence intensity, enzymatic activity, immunoreactivity, DNA hybridization, polymerase chain reaction, etc.

Promiscuous second messenger assays

It is not possible to predict, a priori and based solely upon the GPCR sequence, which of the cell's many different signaling pathways any given orphan receptor will naturally use. It is possible, however, to coax receptors of different functional classes to signal through a pre-selected pathway through the use of promiscuous G_{α} subunits. For example, by providing a cell based receptor assay system with an endogenously supplied promiscuous G_{α} subunit such as $G_{\alpha 15}$ or

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-94-

 $G_{\alpha 16}$ or a chimeric G_{α} subunit such as $G_{\alpha qz}$, a GPCR, which might normally prefer to couple through a specific signaling pathway (e.g., G_s , G_i , G_q , G_0 , etc.), can be made to couple through the pathway defined by the promiscuous G_{α} subunit and upon agonist activation produce the second messenger associated with that subunit's pathway. In the case of $G_{\alpha 15}$, $G_{\alpha 16}$ and/or $G_{\alpha qz}$ this would involve activation of the G_q pathway and production of the second messenger IP3. Through the use of similar strategies and tools, it is possible to bias receptor signaling through pathways producing other second messengers such as Ca^{++} , cAMP, and K^+ currents, for example (Milligan, 1999).

It follows that the promiscuous interaction of the exogenously supplied G_{α} subunit with the orphan receptor alleviates the need to carry out a different assay for each possible signaling pathway and increases the chances of detecting a functional signal upon receptor activation.

20 <u>Methods for recording currents in Xenopus oocytes</u>

Oocytes are harvested from Xenopus laevis and injected with mRNA transcripts as previously described (Quick and Lester, The test receptor of this 1994; Smith et al., 1997). invention and $G\alpha$ subunit RNA transcripts are synthesized using the T7 polymerase ("Message Machine," Ambion) from linearized plasmids or PCR products containing the complete coding region of the genes. Oocytes are injected with 10 ng synthetic receptor RNA and incubated for 3-8 days at 17 Three to eight hours prior to recording, oocytes are injected with 500 pg promiscuous $G\alpha$ subunits mRNA in order to observe coupling to Ca** activated Cl currents. Dual electrode voltage clamp (Axon Instruments Inc.) is performed using 3 M KCl-filled glass microelectrodes having resistances of 1-2 MOhm. Unless otherwise specified, oocytes are voltage clamped at a holding potential of -80 mV.

PCT/US00/12065

-95-

recordings, oocytes are bathed in continuously flowing (1-3 ml/min) medium containing 96 mM NaCl, 2 mM KCl, 1.8 mM CaCl₂, 1 mM MgCl₂, and 5 mM HEPES, pH 7.5 (ND96). Drugs are applied either by local perfusion from a 10 μ l glass capillary tube fixed at a distance of 0.5 mm from the oocyte, or by switching from a series of gravity fed perfusion lines.

Other oocytes may be injected with a mixture of receptor mRNAs and synthetic mRNA encoding the genes for G-protein-activated inward rectifier channels (GIRK1 and GIRK4, U.S. Patent Nos. 5,734,021 and 5,728,535 or GIRK1 and GIRK2) or any other appropriate combinations (see, e.g., Inanobe et al., 1999). Genes encoding G-protein inwardly rectifying K⁺ (GIRK) channels 1, 2 and 4 (GIRK1, GIRK2, and GIRK may be obtained by PCR using the published sequences (Kubb et al., 1993; Dascal et al., 1993; Krapivinsky et al., 1995 and 1995b) to derive appropriate 5' and 3' primers. Human heart or brain cDNA may be used as template together with appropriate primers.

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Heterologous expression of GPCRs in *Xenopus* oocytes has been widely used to determine the identity of signaling pathways activated by agonist stimulation (Gundersen et al., 1983; Takahashi et al., 1987). Activation of the phospholipase C (PLC) pathway is assayed by applying test compound in ND96 solution to oocytes previously injected with mRNA for the mammalian orphan receptor (with or without promiscuous G proteins) and observing inward currents at a holding potential of -80 mV. The appearance of currents that reverse at -25 mV and display other properties of the Ca⁺⁺-activated Cl⁻ (chloride) channel is indicative of mammalian receptoractivation of PLC and release of IP3 and intracellular Ca⁺⁺. Such activity is exhibited by GPCRs that couple to G_{α} or G_{11} .

Measurement of inwardly rectifying K⁺ (potassium) channel

-96-

(GIRK) activity may be monitored in oocytes that have been co-injected with mRNAs encoding the mammalian orphan receptor plus GIRK subunits. GIRK gene products co-assemble to form a G-protein activated potassium channel known to be activated (i.e., stimulated) by a number of GPCRs that couple to G₁ or G₀ (Kubo et al., 1993; Dascal et al., 1993). Oocytes expressing the mammalian orphan receptor plus the GIRK subunits are tested for test compound responsivity by measuring K⁺ currents in elevated K⁺ solution containing 49 mM K⁺.

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In the present invention, oocytes were harvested from Xenopus laevis and injected with mRNA transcripts as previously described (Quick and Lester, 1994; Smith et & .,1997). SNORF36a RNA transcripts were synthesized using the T7 polymerase ("Message Machine", Ambion) from the plasmid BO109 linearized with NotI. Oocytes were injected with 5-25 ng synthetic RNA and incubated for 3-8 days at 17°C. Dual electrode voltage clamp (Axon Instruments Inc.) was performed using 3 M KCl-filled glass microelectrodes having resistances Unless otherwise specified, oocytes were of 1-2 MOhm. voltage clamped at a holding potential of -80 mV. recordings, oocytes were bathed in continuously flowing (1-3 ml/min) medium containing 96 mM NaCl, 2 mM KCl, 1.8 mM CaCl2, 1 mM MgCl₂, 5 mM HEPES, pH 7.5 (ND96), and the appropriate ligand.

Experiments were carried out under minimal light conditions which required compound weighing, final dilutions and oocyte recordings to be performed under photographic darkroom lights ("Brightlab Junior" safelight). In some cases, the ligands were briefly exposed to room light (5-10 min.) during the weighing procedure. For eliciting physiological responses from oocytes, the light stimulus was a 60 W tungsten lamp set at a distance of 30 cm from the oocyte. Drugs were applied

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PCT/US00/12065

-97-

either by superfusion, switching from a series of gravity fed perfusion lines, or by local perfusion from a 10 μ l glass capillary tube fixed at a distance of 0.5 mm from the oocyte. Experiments were carried out at room temperature. All values are expressed as mean \pm standard error of the mean.

Membrane preparations

Cell membranes expressing the receptor protein of this invention are useful for certain types of assays including but not restricted to ligand binding assays, GTP-q-S binding assays, and others. The specifics of preparing such cell membranes may in some cases be determined by the nature of the ensuing assay but typically involve harvesting whole cells and disrupting the cell pellet by sonication cold buffer (e.g. 20 mM Tris HCl, mM EDTA, pH 7.4 at 4° C). The resulting crude cell lysate is cleared of cell debris by low speed centrifugation at 200xg for 5 min at 4° C. The cleared supernatant is then centrifuged at 40,000xq for 20 min at 4° C, and the resulting membrane pellet is washed by suspending in ice cold buffer and repeating the high speed centrifugation step. The final washed membrane pellet is resuspended in assay buffer. Protein concentrations are determined by the method of Bradford (1976) using bovine serum albumin as a standard. The membranes may be used immediately or frozen for later use.

Generation of baculovirus

The coding region of DNA encoding the human receptor disclosed herein may be subcloned into pBlueBacIII into existing restriction sites or sites engineered into sequences 5' and 3' to the coding region of the polypeptides. To generate baculovirus, 0.5 μ g of viral DNA (BaculoGold) and 3 μ g of DNA construct encoding a polypeptide may be cotransfected into 2 x 10^6 Spodoptera frugiperda insect Sf9 cells by the calcium phosphate co-precipitation method, as

-98-

outlined by Pharmingen (in "Baculovirus Expression Vector System: Procedures and Methods Manual"). The cells then are incubated for 5 days at 27°C.

5 The supernatant of the co-transfection plate may be collected by centrifugation and the recombinant virus plaque purified. The procedure to infect cells with virus, to prepare stocks of virus and to titer the virus stocks are as described in Pharmingen's manual.

Localization of mRNA coding for human and rat SNORF36

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<u>Ouantitative RT-PCR using a fluorogenic probe with real time detection</u>: Quantitative RT-PCR using fluorogenic probes and a panel of mRNA extracted from human and total RNA extracted from rat tissue was used to characterize the localization of rat and human SNORF36.

This assay utilizes two oligonucleotides for conventional PCR amplification and a third specific oligonucleotide probe that is labeled with a reporter at the 5' end and a quencher at the 3' end of the oligonucleotide. In the instant invention, FAM (6-carboxyfluorescein) and JOE (6 carboxy-4.5-dichloro-2,7-dimethoxyfluorescein) were the two reporters that were utilized and TAMRA (6-carboxy-4,7,2,7'-tetramethylrhodamine) was the quencher. As amplification progresses, the labeled oligonucleotide probe hybridizes to the gene sequence between the two oligonucleotides used for amplification. activity of thermostable DNA Taq, or rTth nuclease polymerases is utilized to cleave the labeled probe. separates the quencher from the reporter and generates a fluorescent signal that is directly proportional to the amount of amplicon generated. This labeled probe confers a high degree of specificity. Non-specific amplification is not detected as the labeled probe does not hybridize. experiments were conducted in a PE7700 Sequence Detection

PCT/US00/12065

-99-

System (Perkin Elmer, Foster City CA),

<u>Ouantitative RT-PCR</u>: For the detection of RNA encoding SNORF36 receptors, quantitative RT-PCR was performed on mRNA extracted from tissue. Reverse transcription and PCR reactions were carried out in 50 μ l volumes using rTth DNA polymerase (Perkin Elmer).

Primers for human SNORF36 were designed to amplify the long subtype selectively (SNORF36b) or to a region common to both SNORF36a and SNORF36b. Amounts of RNA encoding the short subtype (SNORF36a) were calculated by subtracting the amount of SNORF36b from the amount of total SNORF36. Primers with the following sequences were used:

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Human SNORF36 subtype non-selective:

Forward primer:

SNORF36b2-457F

20 5'-GGCTGCGAGTTCTATGCCTT-3' (SEQ ID NO: 36)

Reverse primer

SNORF36b2-547r

5'-TTACCAGGTAGCGGTCCAGG-3' (SEQ ID NO: 37)

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Fluorogenic oligonucleotide probe:

SNORF36b2-483T

5'(6-FAM)-AGCTCTCTTTGGCATTTCCTCCATGATCA-(TAMRA)3' (SEQ ID NO: 38)

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Human SNORF36b (long subtype) selective:

Forward primer:

SNORF36b b sel-256F

35 5'-CTGGGCAACCTGACGGTC-3' (SEQ ID NO: 39)

-100-

Reverse primer

SNORF36b b sel-346R

5'-CAGGTGTCCGGAGGCTTCT-3' (SEQ ID NO: 40)

Fluorogenic oligonucleotide probe:

SNORF36b b sel-294T

5'(6-FAM)-TGTGCTTCGTGGAGTCACTGTGATGAT-(TAMRA)3' (SEQ ID NO:
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10 Rat SNORF36

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Forward primer SNORF36-rat-34F 5'-TCCACTGGCCACCATCG-3' (SEQ ID NO: 42)

Reverse primer

SNORF36-rat-107R

5'-GGGCATAGAGCCAGACACCTAG-3' (SEQ ID NO: 43)

Fluorogenic oligonucleotide probe:

SNORF36-rat-52T

5'(6-FAM)-CATGAGATCCAAGAGACGGCA-(TAMRA)3' (SEQ ID NO: 44)

Using these primer pairs, amplicon length is 90 bp for human SNORF36-non subtype-selective, 90 bp for human SNORF36b-selective, and 73 bp for rat SNORF36. Each human SNORF36 RT-PCR reaction contained 50 ng mRNA and each rat SNORF36 RT-PCR reaction contained 100 ng total RNA. Oligonuceotide concentrations were: 500 nM of forward and reverse primers, and 200 nM of fluorogenic probe. Concentrations of reagents in each reaction were: 300 µM each of dGTP; dATP; dCTP; 600 µM UTP; 3.0mM Mn(OAc)2; 50 mM Bicine; 115 mM potassium acetate, 8% glycerol, 5 units rTth DNA polymerase, and 0.5 units of uracil N-glycosylase. Buffer for RT-PCR reactions also contained a fluor used as a passive reference (ROX:

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PCT/US00/12065

-101-

Perkin Elmer proprietary passive reference I). All reagents for RT-PCR (except mRNA and oligonucleotide primers) were obtained from Perkin Elmer (Foster City, CA). Reactions were carried using the following thermal cycler profile: 50°C 2 min., 60°C 30 min., 95°C 5 min., followed by 40 cycles of: 94°C, 20 sec., 62°C 1 min.

controls for PCR reactions consisted Positive amplification of the target sequence from a plasmid construct when available. Standard curves for quantitation of human SNORF36 were constructed using the SNORF36 gene in a plasmid construct. RNA extracted from whole brain was used to construct a standard curve for rat SNORF36. Negative controls consisted of mRNA blanks, as well as primer and mRNA blanks. To confirm that the mRNA was not contaminated with genomic DNA, PCR reactions were carried out without reverse transcription using Taq DNA polymerase. Integrity of RNA was assessed by amplification of RNA coding for cyclophilin or glyceraldehyde 3-phosphate dehydrogenase (GAPDH). Following reverse transcription and PCR amplification, analyzed using Perkin Elmer sequence detection software. fluorescent signal from each well was normalized using an internal passive reference, and data was fitted a standard curve to obtain relative quantities of SNORF36 mRNA expression.

Expression of SNORF36 mRNA in human retina .

To determine if SNORF36 mRNA is expressed in retina, RT-PCR was performed on human retina Quick Clone cDNA (Clontech, Palo Alto, CA) using two different primer pairs (BB788, a forward SNORF36 primer, with BB789, a reverse SNORF36 primer; or BB796, a forward SNORF36 primer with BB798, a reverse SNORF36 primer). PCR was performed using the Expand Long Template PCR System (Roche Molecular Biochemicals, Indianapolis, IN) under the following conditions: 94°C hold

-102-

for 5 minutes; 40 cycles of 94°C for 30 seconds, 68°C for 1.5 minutes; 68°C hold for 7 minutes; 4°C hold until ready for agarose gel electrophoresis. As controls, similar PCR reactions were carried out using human whole brain Quick Clone cDNA, human genomic DNA(Clontech, Palo Alto, CA), or water as templates.

BB788 5'- TCCTCCATGATCACCCTGACGGC -3' (SEQ ID NO: 45)

BB789 5'- TCTGGAGAGCCCGTCCTGTCTCC -3' (SEQ ID NO: 46)

BB796 5'- GGCAACCTGACGGTCATCTATACC -3' (SEQ ID NO: 47)

BB798 5'- TTGGACGCCACACCAAAGGTGGCC -3' (SEQ ID NO: 48)

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15 Chromosomal localization for human SNORF36 receptor gene

Chromosomal localization for the human SNORF36 receptor gene was established using a panel of radiation hybrids prepared by the Stanford Human Genome Center (SHGC) and distributed by Research Genetics, Inc. The "Stanford G3" panel of 83 radiation hybrids was analyzed by PCR using the same primers, probes and thermal cycler profiles as used for localization. 20 ng of DNA was used in each PCR reaction. Data was submitted to the RH Server (SHGC) which linked the SNORF36 gene sequences to specific markers. NCBI LocusLink and NCBI GeneMap '99 were used to further analyze the data.

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-103-

RESULTS AND DISCUSSION

Identification of a fragment of the human SNORF36 receptor

A human placental genomic library was screened, under reduced stringency conditions, with oligonucleotide probes directed to the third, fifth and sixth transmembrane regions of the human serotonin 5-HTl_D receptor. Positively-hybridizing clones were isolated, plaque-purified, characterized by Southern blot analysis, and sequenced. One clone, hplOb, contained a 2.3 kb HindIII/XbaI fragment which hybridized with the human 5-HTl_D-derived oligonucleotide probes and was subsequently subcloned into a pUC vector. This clone, called K39, was a partial gene fragment, encoding TMII and MIII and a possible downstream intron of a putative novel GPCR.

Isolation of a full-length human SNORF36 receptor

In order to obtain additional sequence for this receptor, a human hippocampal cDNA library was screened by PCR using primers directed against K39. One positive pool was successfully subdivided until a single clone, TL252, was isolated. DNA sequencing of this clone revealed that it contained TMI through TMVII but was lacking the NH_2 and COOH termini. The remaining sequence, including the initiating methionine and the stop codon were obtained by 5' and 3' RACE performed on human hippocampal cDNA. The human hippocampal cDNA library was re-screened and two positive clones were 260-13-1 and 143-33-3, which contained TMI identified, These two clones both had an 11 through the stop codon. amino acid insert in the first intracellular loop that was not present in TL252.

A full-length cDNA clone for SNORF36a (without the 11 amino acid insert) was obtained by ligating together a 1021 bp <code>BamHI/BglI</code> fragment from a hippocampal cDNA PCR product along

WO 00/66630 PCT/US00/12065

-104-

with a 1180 bp BglI/EcoRI fragment from the human hippocampal library pool 260.13.1 into a BamHI/EcoRI-cut pcDNA3.1. This construct, B0108, had a single nucleotide mutation which was corrected by site-directed mutagenesis. The corrected construct, BO109, was renamed pcDNA3.1-hSNORF36a-f. Analysis of 9 independent PCR products from human hippocampal cDNA revealed the presence of an allelic variation. 39 (Figures 1A-1C and 3A-3C) is either an adenine or a guanine. A full-length clone for SNORF36b (with the 11 amino acid insert) was obtained by ligating together a 1130 bp NspI/HindIII fragment from a PCR of B0108 along with a 390 bp BamHI/NspI fragment from one of the PCR products described above into a BamHI/HindIII-cut pcDNA3.1. The resulting SNORF36b construct, B0110, was renamed pcDNA3.1-hSN 'RF36b-f.

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in SNORF36a is 1434 open reading frame largest The nucleotides, which is predicted to encode a protein of 478 Using a downstream methionine results in an amino acids. open frame of 1377 nucleotides and is predicted to encode a protein of 459 amino acids. The nucleotide and amino acid sequences of SNORF36a are shown in Figures 1A-1C and 2A-2B, respectively. The largest open reading frame in SNORF36b is 1467 nucleotides, which is predicted to encode a protein of 489 amino acids. Using a downstream methionine results in an open frame of 1410 nucleotides and is predicted to encode a protein of 470 amino acids. The nucleotide and amino acid sequences of SNORF36b are shown in Figures 3A-3C and 4A-4C Hydropathy analysis of both SNORF36a and respectively. SNORF36b protein are consistent with a putative topography of seven transmembrane domains, indicative of the G proteincoupled receptor family.

A comparison of nucleotide and peptide sequences of human SNORF36a and SNORF36b with sequences contained in the Genbank, EMBL, and SwissProtPlus databases reveals that the

WO 00/66630

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PCT/US00/12065

-105-

amino acid sequences of these clones are most related to the Xenopus melanopsin receptor (45% identity), giant octopus rhodopsin (38% identity), giant scallop rhodopsin (36% identity), Japanese flying squid and cuttlefish rhodopsins (35% identities), squid rhodopsin (34% identity), human peropsin and human RGR (31% identities), human encephalopsin (30% identity), human rhodopsin (28% identity) and bovine rhodopsin (27% identity). There were no sequences in the Genbank databases (Genembl, STS, EST, GSS, or SwissProt) that were identical to SNORF36.

Human SNORF36a and SNORF36b have seven potential protein kinase C (PKC) phosphorylation motifs at serine 111 in the first intracellular loop (numbers in this sec ion are relative to SNORF36b in Figures 4A-4C), serine 194 in the second intracellular loop, threonine 276 in the third intracellular loop, and at serines 386, 395, and 415 and threonine 399 in the carboxy-terminal tail. There is also one potential N-linked glycosylation site at asparagine 88 in the first transmembrane domain and one cAMP phosphorylation site at serine 390. SNORF36a and SNORF36b also have two potential casein kinase II phosphorylation sites at serine 425 and threonine 432 in the carboxy-terminal tail.

25 <u>Isolation of a Fragment of the Rat Homologue of SNORF36</u>

A fragment of the rat homologue of SNORF36 was amplified from rat genomic DNA by low stringency PCR using oligonucleotide primers designed against the human SNORF36. The sequence of this fragment was then used to generate rat SNORF36 PCR primers which were used to amplify under high stringency a SNORF36 fragment from rat spinal cord cDNA. This fragment, KO56, contains 250 nucleotides of rat SNORF36, from the 3' end of TMII to the middle of TMV. The nucleotide and amino acid sequences of the rat SNORF36 fragment are shown in Figures 5 and 6, respectively. The rat SNORF36 fragment

WO 00/66630

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PCT/US00/12065

-106-

shares 86% nucleotide and 88% amino acid identities with the human SNORF36 (Figures 7 and 8). KO56 was renamed pEXJ-T3T7-rSNORF36p. There were no sequences in the Genbank databases (Genembl, STS, EST, GSS, or SwissProt) that were identical to rat SNORF36.

Isolation of a full-length rat SNORF36 receptor

In order to obtain a full-length cDNA for rat SNORF36, a rat spinal cord cDNA library was screened by PCR using primers directed against the rat SNORF36 fragment, KO56. One positive pool was successfully subdivided until a single clone, BO132, was isolated. DNA sequencing of this clone revealed that it contained an open reading frame of 1422 nucleotides, 148 nucleotides of 5'UT and approximately 1000 nucleotides of 3'UT. The construct BO132, in the expression vector pEXJ.T7, was renamed pEXJ.T7-rSNORF36-f.

The largest open reading frame in the rat SNORF36 construct BO132 is 1422 nucleotides, which is predicted to encode a protein of 474 amino acids. The nucleotide and amino acid sequences of rat SNORF36 are shown in Figures 9A-9C and 10A-10C, respectively. Hydropathy analysis of the rat SNORF36 protein are consistent with a putative topography of seven transmembrane domains, indicative of the G protein-coupled receptor family.

Rat SNORF36 shares 81% nucleotide identity and 79% amino acid identity with human SNORF36a (Figures 11A-11D and 12A-12B, respectively). A comparison of nucleotide and peptide sequences of rat SNORF36 with sequences contained in the Genbank, EMBL, and SwissProtPlus databases reveals that the amino acid sequences of these clones are most related to the Xenopus melanopsin receptor (46% identity), giant octopus rhodopsin and giant scallop rhodopsin (35% identities), cuttlefish rhodopsin (34% identity), Japanese flying squid

WO 00/66630 PCT/US00/12065

-107-

and mouse peropsin (33% identities), human RGR (32% identity), human and bovine rhodopsin (28% identities) and human encephalopsin (27% identity). There were no sequences in the Genbank databases (Genembl, STS, EST, GSS, or SwissProt) that were identical to rat SNORF36.

SNORF36 has six potential protein kinase C (PKC) Rat phosphorylation motifs at serine 183 in the second intracellular loop, threonine 265 in the third intracellular loop, and at serines 381 and 385, and threonines 460 and 463 in the carboxy-terminal tail. There are also three potential N-linked glycosylation sites at asparagines 31 and 35 in the amino terminal tail and at asparagine 88 in the first transmembrane domain. There is one cAMP phosphorylation site at threonine 187 in the fourth transmembrane domain, and three potential casein kinase II phosphorylation sites at serines 411 and 452 and threonine 418 in the carboxy-terminal tail.

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20 <u>Ca²⁺ Mobilization and Phosphoinositide Hydrolysis in</u> SNORF36a-expressing Cells

Since SNORF36a is similar to invertebrate opsins, it was hypothesized that it may couple to G_q G-protein and may induce phosphoinositide hydrolysis followed by release of intracellular Ca^{2+} . Therefore, Ca^{2+} mobilization and inositol phosphate (IP) accumulation were evaluated in hSNORF36a-transfected Cos-7 cells. Exposure to laser light of hSNORF36a-transfected Cos-7 cells, kept in dark, in the FLIPR resulted in Ca^{2+} mobilization in the absence of any exogenous ligand (Figure 13A). This observation suggested that an endogenous chromophore, probably a retinal isomer, was already attached to the hSNORF36a receptor, and exposure to laser light resulted in photoisomerization of the ligand and activation of the receptor. Since this situation did not

WO 00/66630 PCT/US00/12065

-108-

allow us to study the effect of an exogenously added ligand, we 'photobleached' the hSNORF36a- and vector-transfected Cos-7 cells by exposing them to light for 90-120 minutes prior Photobleaching is a phenomenon where to the experiment. exposure to light results in photoisomerization of endogenous ligand, eventually leading to release of the ligand and formation of an opsin with an empty binding This technique is used widely to dissociate the endogenous ligand from rhodopsin. Upon photobleaching, the Ca2+ mobilization response to laser light in the hSNORF36atransfected Cos-7 cells was either reduced or completely implying dissociation (Figure 13B), abolished Therefore, photobleaching was used for endogenous ligand. all subsequent experiments unless indicated otherwise.

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Opsins can be activated by several retinal analogues, including all-trans-retinal (Surya and Knox, 1998) which is a 'natural' agonist formed upon photoisomerization of 11-cis-Upon addition of retinal analogues to photobleached Ca²⁺ SNORF36a-transfected Cos-7 cells, significant mobilization was observed (Figures 14A, 15A). The rank order of potency was 9-cis-retinal (EC₅₀:152nM) > all-trans-retinal $(EC_{50}:263 \text{ nM}) > 13-cis\text{-retinal } (EC_{50}:477\text{nM}) \text{ (See Figure 15A)}.$ similar response to all-trans-retinal was photobleached SNORF36a-transfected HEK293 cells (Figure 15B). In contrast, retinals did not change intracellular Ca2+ levels in the vector-transfected Cos-7 cells (Figure 14B). Collectively, these results indicate that SNORF36a is an opsin which can be activated by retinals. To confirm these results further, we examined the effect of SNORF36a on IP Concentration-dependent increase accumulation was seen upon all-trans-retinal addition (Figure 16A) in SNORF36a-transfected Cos-7 cells exposed to light but not in vector-transfected cells (Figure 16A). In contrast, in SNORF36a-transfected Cos-7 cells not exposed to light,

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-109-

all-trans-retinal did not change 3H-IP levels, indicating that light may be required either for dissociation of endogenous ligand or activation of the receptor (Figure 16B). Collectively, these results suggest that activation of hSNORF36a can induce phosphoinositide hydrolysis which may be upstream of intracellular Ca2+ mobilization. Finally, we tested the potential antagonism by β -ionone which is to retinals. structurally similar β-ionone, concentration, significantly antagonized responses to alltrans-retinal and 9-cis-retinal in hSNORF36a-transfected Cos-7 cells (Figures 17A and 17B). The antagonism by β -ionone appeared to be insurmountable (Figures 17A and 17B). may be due to the intrinsic characteristics of β -iononehSNORF36a interaction. Alternatively, it may represent the hemi-equilibrium conditions achieved in the Nonetheless, these results confirm that retinal-like structure can bind to this receptor. These results are consistent with the notion that SNORF36a is an opsin coupled to Ca2+ mobilization and phosphoinositide hydrolysis, most likely via G_o G-protein, and is activated by retinals.

Photic Responses From Oocytes Expressing SNORF36a

The experiments with mammalian cells suggest that SNORF36a, stimulated by the presence of retinoids, can activate a second messenger response that includes release of inositol triphosphates and intracellular free Ca**. Heterologous expression in Xenopus oocytes has been widely used to study the ligand binding and functional activity of a variety of G-protein coupled receptors (Gundersen et al., Takahashi et al., 1987), including rhodopsin (Khorana et al., 1988). Activation of the PLC pathway in oocytes leads to stimulation of the endogenous Ca⁺⁺-activated Cl⁻ current. sought to determine if oocytes expressing SNORF36a could be stimulated by retinoids, and if sensitivity to retinoids was dependent upon activation by light.

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-110-

Upon acute administration of retinoids to occytes expressing SNORF36a, no ionic currents were detected under dark conditions or subsequent to the light stimulus. This result suggested that preincubation of ligands may be necessary to see activation of the receptors. To detect light-dependent ionic responses in Xenopus oocytes expressing bovine opsin, Khorana and co-workers (1988) incubated oocytes for 30-40 minutes with 11-cis retinal to permit the formation of rhodopsin. In another example, the photosensitivity of Xenopus melanophores has been seen to occur only following for several hours preincubation with all-trans retinal (Rollag, 1996). Following these examples, oocytes expressing SNORF36a were pre-incubated in the dark for 24-48 h with alltrans-retinal, 13-cis-retinal, or all-trans-retingic acid (ATRA) (each at 100 nM) in $ND96^{++}$ (Figure 19). preincubated with any of these ligands, the oocytes produced rapidly desensitizing Cl currents in response to the light stimulus (Figures 18A and 19). Current amplitudes ranged from 100 to 500 nA, and appeared quite similar in waveform to those elicited by stimulation of receptors that are known to activate PLC (Gundersen et al., 1983). The currents generated by SNORF36a stimulation were considerably larger in amplitude than those reported for oocytes expressing bovine rod opsin (about 5 nA, Khorana et al., 1988). reason for this is likely due to the different signal transduction pathways activated by these two receptors. SNORF36a may evoke release of intracellular activation of $G\alpha_{\alpha}$, whereas vertebrate opsin couples to transducin and cannot directly activate the PLC pathway. Except for ATRA, the retinoids that were exposed to room light before incubation with the oocytes expressing SNORF36a produced significantly larger current amplitudes (about 5fold) than non-exposed retinoids (Figures 18B and 19). Control oocytes, not injected with SNORF36a mRNA, that were pre-incubated with all-trans-retinal or 13-cis-retinal did

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-111-

not respond to light (Figure 18C; n = 10 oocytes).

Detection of RNA coding for human SNORF36 receptors

mRNA was isolated from multiple tissues (listed in Table 1) and assayed as described.

Human SNORF36

Quantitative RT-PCR using a fluorgenic probe demonstrated mRNA encoding human SNORF36 to be localized in highest abundance in CNS tissue. All CNS tissues assayed demonstrate measurable levels of SNORF36 RNA.

Highest levels are found in the caudate-putamen, amygdala and hippocampus. Localization of high levels of SNORF36 RNA to the caudate-putamen indicates a role in modulation of dopaminergic function, or the modulation of extrapyramidal motor systems. High levels of SNORF36 RNA in the hippocampal formation and amygdala support the hypothesis that SNORF36 is involved in the modulation of learning and memory. also have a role in the regulation of fear, mood, and may provide a target for the treatment of depression, anxiety, phobias and mood disorders. Other regions of the CNS containing SNORF36 RNA include the spinal cord and thalamus, implying an important role in sensory transmission or modulation (including nociception). SNORF36 is also expressed in lower levels in the substania nigra, hypothalamus, and cerebellum. The broad distribution of SNORF36 RNA throughout the CNS implies a modulatory role in multiple systems within the CNS.

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Fetal brain, although expressing SNORF36 mRNA does so in much lower levels than that found in the adult. In fetal brain, SNORF36 RNA is barely detectable. There is a 70-fold difference in mRNA levels between fetal and adult brain. It is not known at this time if the developmental regulation

WO 00/66630 PCT/US00/12065

-112-

occurs in all regions within the CNS or is restricted to selected areas. The time course of this increase has not been examined and would be important in understanding the function of this receptor.

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In peripheral tissue, most tissues assayed expressed measurable SNORF36 mRNA levels. The peripheral tissues expressing the highest levels of SNORF36 mRNA are skeletal muscle, heart and small intestine. Levels in these tissues are 10-fold lower than that detected in the highest CNS regions. Presence of SNORF36 mRNA in these areas implies a role in regulation of contractility, perhaps by a common mechanism. Other tissues assayed contain low levels of SNORF36 mRNA as indicated in Table 1.

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Both subtypes (SNORF36a and SNORF36b) were assayed in all tissue samples. Primers for human SNORF36 were designed to amplify the long subtype selectively (SNORF36b) or to a region common to both SNORF36a and SNORF36b. Amounts of RNA encoding the short subtype (SNORF36a) were calculated by subtracting the amount of SNORF36b from the amount of total SNORF36. In most regions assayed, SNORF36a is the predominant subtype expressed. The exceptions are the caudate-putamen and fetal lung. In these tissues, SNORF36a accounts for 31% and 43%, respectively, of the total SNORF36. However, in all other tissue assayed, SNORF36a accounts for 60% to 100% of the total SNORF36 present in tissue.

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In summary, the distribution of human SNORF36 mRNA implies broad regulatory function in the CNS, most notably in modulation of extrapyramidal motor systems, modulation of the limbic system, and sensory transmission. Its presence, albeit at lower levels, in peripheral tissues implies a broad regulatory role in multiple organ systems. The predominance of SNORF36a in most tissue indicates that functionally, it is the dominant subtype.

WO 00/66630

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PCT/US00/12065

-113-

Rat SNORF36

As with the human SNORF36 receptor mRNA, all central nervous system structures assayed contain rat SNORF36 RNA. However, the highest levels of rat SNORF36 RNA are found in the retina The retina contains 5-fold more RNA encoding SNORF36 than any other tissue. The high levels found in the retina imply a function in vision or photoentrainment. The hypothesis that rat SNORF36 plays a role in circadian rhythm is supported by the fact that substantial levels of SNORF36 RNA are also expressed in the pineal gland, as well as the hypothalamus (Table 2). The localization and functional data imply that SNORF36 is a non-rod, non-cone, ocular photoreceptor. It has been suggested that non-rod, non-cone ocular photoreceptors are responsible for photoentrainment and regulation of melatonin synthesis in the pineal gland (Freedman et al., 1999, Lucas et al., 1999).

Other regions containing high levels of SNORF36 RNA include trigeminal ganglia, spinal cord, and medulla. Localization to these structures as well as moderate levels in the dorsal root ganglia, strongly suggest a role in sensory transmission (or modulation) including nociceptive stimuli. Localization in the medulla also implies regulation of autonomic centers controlling respiration and cardiovascular function. CNS regions containing high levels of SNORF36 RNA include the amygdala, and substanta nigra. High levels in the amygdala and other limbic (or limbic related) structures suggests a role in modulation of mood, fear, phobia, anxiety and may provide a therapeutic target for the treatment of depression and other neuropsychiatric disorders. Localization to the substantia nigra (in conjunction with expression in the striatum) implies a role in regulation of dopaminergic systems, and may provide a therapeutic target for treatment of movement disorders such as Parkinsons disease or tardive dyskinesea.

WO 00/66630 PCT/US00/12065

-114-

The high levels of SNORF36 mRNA expressed in the hypothalamus described previously, in addition to the potential role in regulation of circadian rhythm, also indicates a role in neuroendocrine regulation, regulation of appetite and other functions that are modulated by the hypothalamus. High levels in the amygdala suggest a role in modulation of mood, fear, phobia, anxiety and may provide a therapeutic target for the treatment of depression and other neuropsychiatric disorders.

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The presence of lower levels of SNORF36 RNA in other areas such as the hippocampal formation, olfactory bulb, cerebral cortex, cerebellum and other areas suggests multiple diverse functions as suggested in Table 2.

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Non-neuronal tissue expressing high levels of SNORF36 RNA include the ovary and testes. Levels in these areas are comparable to many CNS regions including the cerebral cortex and cerebellum (Table 2). This strongly suggests a role in endocrine regulation or reproductive function, by neuronal or extraneuronal mechanisms. Other peripheral tissues showing moderate amounts of SNORF36 mRNA are listed in Table 2.

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In summary, the high levels or SNORF36 RNA in the retina, pineal gland, and hypothalamus suggest a role as a non-rod, non-cone ocular photoreceptor involved in regulation of circadian rhythms. The distribution of rat SNORF36 mRNA throughout the CNS implies broad regulatory function in the nervous system. Potential functions include modulation of sensory transmission, modulation of extrapyramidal motor systems, and modulation of the limbic system. The ovary and testes are among multiple peripheral organs that may be regulated by SNORF36. Its presence, albeit at lower levels, in other peripheral tissues implies a broad regulatory role in multiple organ systems.

-115-

Table 1

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Summary of distribution of mRNA coding for human SNORF36 receptors (long and short forms).

Amounts of mRNA encoding human SNORF36 expressed as, mean (whole brain standard curve), % of highest expressing tissue (CPu \pm SEM).

SNORF36 SNORF SNORF SNORF Potential Region % of 36b 36a 36a % applica-% of % of of max tions max SNORF36 max 70.78 amygdala 50.10 14.64 35.47 Depression ±5.73 phobias, anxiety, mood disorders 100.00 69.18 30.82 30.82 Modulation caudate-+7.89 of putamen dopaminergic function, Modulation of extrapyramidal motor systems 0.31 cerebellum 0.31 not 100 Motor detecte ±0.07 coordination 0.22 0.01 0.22 97.28 Developfetal ±0.05 mental brain disorders

-116-

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Region	SNORF36	SNORF	SNORF	SNORF	Potential
	% of	365	36a	36a &	applica-
}	max	% of	₹ of	of	tions
Ì		max	max	SNORF36	
fetal	0.61	not	0.61	100	Develop-
kidney	<u>+</u> 0.10	detecte			mental
		d			disorders
fetal	0.04	0.01	0.03	63.08	Develop-
liver	±0.01		i		mental
					disorders
fetal lung	0.42	0.24	0.18	42.79	Develop-
	±0.11				mental
					disorders
heart	7.44	not	7.44	100	Cardio-
	<u>+</u> 2.46	detecte			vascular
		d			disorders
hippo-	26.28	10.52	15.76	59.96	Cognition/
campus	±1.76				memory
hypo-	0.77	0.04	0.72	94.49	appetite/
thalamus	<u>+</u> 0.07				obesity,
			·		neuro-
					endocrine
					regulation
kidney	0.16	trace	0.16	100	Hyperten-
	<u>+</u> 0.04				sion,
		i			electro-
					lyte
					balance
liver	trace	trace	0.02	NA	Diabetes
lung	trace	trace	0.03	NА	Respira-
			ļ		tory
		ļ	ļ		disorders,
					asthma
pancreas	0.07	not	0.07	100	Diabetes,
	±0.01	detecte			endocrine
		d			disorders

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PCT/US00/12065

-117-

Region	SNORF36	SNORF	SNORF	SNORF	Potential
	% of	366	36a	36a %	applica-
	max	% of	to f	o£	tions
		max	max	SNORF36	
pituitary	4.38	0.17	4.21	96.20	Endocrine/
	<u>+</u> 0.78				neuro-
	ł				endocrine
:	Į.				regulation
placenta	0.03	0.05	0.03	100	Gestation-
	<u>+</u> 0.01	<u>+</u> 0.02			al
			1		abnormali-
·					ties
small	6.24	0.34	5.90	94.51	Gastro-
intestine	±1.51				intestinal
					disorders
spinal	7.60	0.06	7.54	99.18	Analgesia,
cord	<u>+</u> 1.38				sensory
					modulation
					and trans-
					mission
spleen	0.10	trace	0.10	100	Immune
	<u>+</u> 0.02	+ =====	trace	NA	disorders Gastro-
stomach	trace	trace	trace	INA	intestinal
	10.51	0.30	10.21	97.15	disorders Musculo-
skeletal	1	0.30	10.21	3,.10	skeletal
muscle	<u>+</u> 0.88		ŀ	-	disorders
substantia	3.73	0.17	3.57	95.48	Modulation
	<u>+</u> 0.22				of
nigra	-0.22				dopamine-
			1		rgic
					function.
					Modulation
		1			
}			1		of motor
					coordina-
l		1	ļ	}	tion.

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-118-

Region	SNORF36	SNORF	SNORF	SNORF	Potential
	% of	36b	36a	36a %	applica-
	max	% of	% of	of	tions
		max	max	SNORF36	
thalamus	5.56	1.19	4.37	78.62	Sensory
	<u>+</u> 0.70				integra-
					tion
		<u> </u>			disorders
whole	15.82	4.97	10.85	68.59	
brain	<u>+</u> 2.28			<u> </u>	

-119-

Table 2

Summary of distribution of mRNA coding for rat SNORF36 receptors

5 mRNA encoding SNORF36 is expressed as % of highest expressing tissue: retina ± SEM.

Tissue	SNORF36	Potential applications
adipose	1.93	metabolic disorders
	<u>+</u> 0.21	
adrenal cortex	2.62	regulation of steroid
	±0.48	hormones
adrenal medulla	2.86	regulation of epinephrine
	±0.24	release
amygdala	9.52	depression, phobias,
	<u>±</u> 0.75	anxiety, mood disorders
aorta	1.08	cardiovascular disorders
	±0.07	
celiac plexus	1.58	modulation of autonomic
	±0.14 4.81	function
cerebellum	4.81	motor coordination
	±0.43 6.35	
cerebral cortex	6.35	Sensory and motor
	±0.41 7.02	integration, cognition
choroid plexus	7.02	regulation of
	±0.93	cerebrospinal fluid
colon	0.61	gastrointestinal
	±0.05 4.18	disorders
dorsal root	4.18	sensory transmission
ganglia	<u>+</u> 0.23	
duodenum	0.86	gastrointestinal
	±0.10	disorders
heart	1.79	cardiovascular
	<u>+</u> 0.19	indications
hippocampus	±0.19 3.45	cognition/memory
	<u>±</u> 0.37	
hypothalamus	9.13	appetite/obesity,
	±0.62	neuroendocrine regulation

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-120-

Tissue	SNORF36	Potential applications
kidney	0.95	electrolyte balance,
•	<u>+</u> 0.07	hypertension
liver	trace	diabetes
medulla	9.33	analgesia, motor
	<u>+</u> 0.57	coordination
nucleus accumbens	4.29	regulation of
	<u>+</u> 0.36	dopaminergic function,
		drug addiction,
		neuropsychiatric
		disorders
olfactory bulb	3.96	olfaction
-	+0.33	
ovary	<u>+</u> 0.33 6.83	reproductive function
	+0.66	
pancreas	<u>+</u> 0.66 0.30	diabetes, endocrine
•	±0.04 8.37	disorders
pineal	8.37	regulation of melatonin
	<u>+</u> 0.21	release
pituitary	±0.21 3.68	endocrine/neuroendocrine
	<u>+</u> 0.37	regulation
retina	±0.37	visual disorders,
	<u>+</u> 7.26	circadian rhythms
salivary gland	<u>+</u> 7.26 1.26	
	±0.21	1
spinal cord	±0.21 14.47	analgesia, sensory
	<u>+</u> 0.44	modulation and
		transmission
spleen	0.27	immune disorders
_	±0.14	
stomach	0.30	gastrointestinal
	<u>+</u> 0.02	disorders
striated muscle	2.08	musculoskeletal disorders
	<u>+</u> 0.18	
striatum	3.72	modulation of
	<u>+</u> 0.67	dopaminergic function,
		motor disorders

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-121-

	SNORF36	Potential applications
Tissue	11.92	modulation of
substantia nigra		
	±2.46	dopaminergic function,
		modulation of motor
		coordination
testes	5.48	reproductive function
	<u>+</u> 0.21	
thalamus	3.85	sensory integration
	<u>+</u> 0.16	disorders
thymus	3.07	immune disorders
	<u>+</u> 0.37	
trigeminal	19.13	sensory transmission
ganglia	<u>+</u> 2.83	
urinary bladder	1.48	urinary incontinence
	<u>+</u> 0.11	
uterus	1.16	reproductive disorders
-	±0.14	
vas deferens	2.85	reproductive function
	<u>+</u> 0.21	
whole brain	4.12	
	±0.39	

Expression of SNORF36 mRNA in human retina

To determine if SNORF36 mRNA is expressed in retina, RT-PCR was performed in human retina cDNA using two pairs of SNORF36-specific primers. Single bands of the expected sizes were visualized following agarose gel electrophoresis for each primer pair. Somewhat weaker bands were observed in similar RT-PCR reactions using whole brain cDNA as the template. In contrast, no bands were observed in RT-PCR reactions in which water was used as the template. As both primer pairs were designed to cross introns, similar sized bands were not detected in RT-PCR reactions in which genomic DNA was used as the template. These results demonstrate that SNORF36 mRNA is expressed in human retina.

WO 00/66630

PCT/US00/12065

-122-

Chromosomal localization of human SNORF36 receptor genes

The human SNORF36 gene maps to SHGC-57810 and SHGC-57490 which is localized to chromosome 10q22.3.

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PCT/US00/12065

-123-

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-132-

What is claimed is:

- An isolated nucleic acid encoding a mammalian SNORF36 receptor.
- 2. The nucleic acid of claim 1, wherein the nucleic acid is DNA.
- 3. The DNA of claim 2, wherein the DNA is cDNA.
- 4. The DNA of claim 2, wherein the DNA is genomic DNA.
 - 5. The nucleic acid of claim 1, wherein the nucleic acid is RNA.
- 6. The nucleic acid of claim 1, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor.
- 7. The nucleic acid of claim 1, wherein the mammalian SNORF36 receptor is a human SNORF36b receptor.
 - 8. The nucleic acid of claim 6, wherein the human SNORF36a receptor has an amino acid sequence identical to that encoded by the plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. PTA-203977).
 - 9. The nucleic acid of claim 7, wherein the human SNORF36b receptor has an amino acid sequence identical to that encoded by the plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. PTA-203976).
 - 10. The nucleic acid of claim 1, wherein the mammalian SNORF36 receptor is a rat SNORF36 receptor.
- 35 11. The nucleic acid of claim 10, wherein the rat SNORF36 receptor has an amino acid sequence identical to that

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-133-

encoded by the plasmid pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216).

- 12. The nucleic acid of claim 6, wherein the human SNORF36a receptor has an amino acid sequence identical to the amino acid sequence shown in Figures 2A-2B (SEQ ID NO: 2).
- 13. The nucleic acid of claim 7, wherein the human SNORF36b receptor has an amino acid sequence identical to the amino acid sequence shown in Figures 4A-4C (SEQ ID NO: 4).
- 14. The nucleic acid of claim 10, wherein the rat SNORF36 receptor has an amino acid sequence identical to the amino acid sequence shown in Figures 10A-10C (SEQ ID NO: 8).
 - 15. A purified mammalian SNORF36 receptor protein.
- 16. The purified mammalian SNORF36 receptor protein of claim 15, wherein the SNORF36 receptor protein is a human SNORF36a receptor protein.
- 25 17. The purified mammalian SNORF36 receptor protein of claim 15, wherein the SNORF36 receptor protein is a human SNORF36b receptor protein.
- 18. The purified mammalian SNORF36 receptor protein of claim 15, wherein the SNORF36 receptor protein is a rat SNORF36 receptor protein.
 - 19. A vector comprising the nucleic acid of claim 1.
- 35 20. A vector comprising the nucleic acid of claim 6 or claim 7.

-134-

- 21. A vector comprising the nucleic acid of claim 10.
- 22. A vector of claim 19, 20 or 21 adapted for expression in a cell which comprises the regulatory elements necessary for expression of the nucleic acid in the cell operatively linked to the nucleic acid encoding the receptor so as to permit expression thereof, wherein the cell is a bacterial, amphibian, yeast, insect or mammalian cell.

23. The vector of claim 22, wherein the vector is a baculovirus.

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- 24. The vector of claim 19, wherein the vector is a plasmid.
 - 25. The plasmid of claim 24 designated pcDNA3.1-hSNORF36a-f (ATCC Accession No. PTA-203977)
- 26. The plasmid of claim 24 designated pcDNA3.1-hSNORF36b-f (ATCC Accession No. PTA-203976).
 - 27. The plasmid of claim 24 designated pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216).
 - 28. A cell comprising the vector of claim 22.
 - 29. A cell of claim 28, wherein the cell is a non-mammalian cell.
 - 30. A cell of claim 29, wherein the non-mammalian cell is a Xenopus oocyte cell or a Xenopus melanophore cell.
- 31. A cell of claim 28, wherein the cell is a mammalian celi.

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-135-

- 32. A mammalian cell of claim 31, wherein the cell is a COS-7 cell, a 293 human embryonic kidney cell, a NIH-3T3 cell, a LM(tk-) cell, a mouse Yl cell, or a CHO cell.
- 33. A cell of claim 28, wherein the cell is an insect cell.
- 34. An insect cell of claim 33, wherein the insect cell is an Sf9 cell, an Sf21 cell or a Trichoplusia ni 5B-4 cell.
- 35. A membrane preparation isolated from the cell of any one of claims 28, 29, 31, 32, 33 or 34.
- 15 36. A nucleic acid probe comprising at least 15 nucleotides, which probe specifically hybridizes with a nucleic acid encoding a mammalian SNORF36 receptor, wherein the probe has a sequence complementary to a unique sequence present within one of the two strands of the nucleic acid encoding the human SNORF36a receptor contained in plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. PTA-203977).
- acid probe comprising at least 15 37. nucleotides, which probe specifically hybridizes with 25 a nucleic acid encoding a mammalian SNORF36 receptor, wherein the probe has a sequence complementary to a unique sequence present within one of the two strands of the nucleic acid encoding the human SNORF36b 30 receptor contained in plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. PTA-203976).
- 38. A nucleic acid probe comprising at least 15 nucleotides, which probe specifically hybridizes with a nucleic acid encoding a mammalian SNORF36 receptor, wherein the probe has a sequence complementary to a

-136-

unique sequence present within one of the two strands of the nucleic acid encoding the mammalian SNORF36 receptor contained in plasmid pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216).

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39. A nucleic acid probe comprising at least 15 nucleotides, which probe specifically hybridizes with a nucleic acid encoding a mammalian SNORF36 receptor, wherein the probe has a sequence complementary to a unique sequence present within (a) the nucleic acid sequence shown in Figures 1A-1C (SEQ ID NO: 1) or (b) the reverse complement thereof.

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40. A nucleic acid probe comprising at least 15 nucleotides, which probe specifically hybridizes with a nucleic acid encoding a mammalian SNORF36 receptor, wherein the probe has a sequence complementary to a unique sequence present within (a) the nucleic acid sequence shown in Figures 3A-3C (SEQ ID NO: 3) or (b) the reverse complement thereof.

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41. A nucleic acid probe comprising at least 15 nucleotides, which probe specifically hybridizes with a nucleic acid encoding a mammalian SNORF36 receptor, wherein the probe has a sequence complementary to a unique sequence present within (a) the nucleic acid sequence shown in Figures 9A-9C (SEQ ID NO: 7) or (b) the reverse complement thereof.

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42. The nucleic acid probe of claim 39, 40 or 41, wherein the nucleic acid is DNA.

43. The nucleic acid probe of claim 39, 40 or 41, wherein the nucleic acid is RNA.

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44. An antisense oligonucleotide having a sequence capable

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-137-

of specifically hybridizing to the RNA of claim 5, so as to prevent translation of the RNA.

- 45. An antisense oligonucleotide having a sequence capable of specifically hybridizing to the genomic DNA of claim 4, so as to prevent transcription of the genomic DNA.
- 46. An antisense oligonucleotide of claim 44 or 45, wherein the oligonucleotide comprises chemically modified nucleotides or nucleotide analogues.
 - 47. An antibody capable of binding to a mammalian SNORF36 receptor encoded by the nucleic acid of claim 1.
- 48. An antibody of claim 47, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor, a human SNORF36b receptor or a rat SNORF36 receptor.
- 49. An agent capable of competitively inhibiting the binding of the antibody of claim 47 to a mammalian SNORF36 receptor.
 - 50. An antibody of claim 47, wherein the antibody is a monoclonal antibody or antisera.
 - 51. A pharmaceutical composition comprising (a) an amount of the oligonucleotide of claim 44 capable of passing through a cell membrane and effective to reduce expression of a mammalian SNORF36 receptor and (b) a pharmaceutically acceptable carrier capable of passing through the cell membrane.
 - 52. A pharmaceutical composition of claim 51, wherein the oligonucleotide is coupled to a substance which inactivates mRNA.

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-138-

- 53. A pharmaceutical composition of claim 52, wherein the substance which inactivates mRNA is a ribozyme.
- 54. A pharmaceutical composition of claim 52, wherein the pharmaceutically acceptable carrier comprises a structure which binds to a mammalian SNORF36 receptor on a cell capable of being taken up by the cells after binding to the structure.
- 10 55. A pharmaceutical composition of claim 54, wherein the pharmaceutically acceptable carrier is capable of binding to a mammalian SNORF36 receptor which is specific for a selected cell type.
- 56. A pharmaceutical composition which comprises an amount of the antibody of claim 47 effective to block binding of a ligand to a human SNORF36 receptor and a pharmaceutically acceptable carrier.
- 57. A transgenic, nonhuman mammal expressing DNA encoding a mammalian SNORF36 receptor of claim 1.
 - 58. A transgenic, nonhuman mammal comprising a homologous recombination knockout of the native mammalian SNORF36 receptor.
- 59. A transgenic, nonhuman mammal whose genome comprises antisense DNA complementary to the DNA encoding a mammalian SNORF36 receptor of claim 1 so placed within the genome as to be transcribed into antisense mRNA which is complementary to and hybridizes with mRNA encoding the mammalian SNORF36 receptor so as to thereby reduce translation of such mRNA and expression of such receptor.
 - 60. The transgenic, nonhuman mammal of claim 57 or 58,

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-139-

wherein the DNA encoding the mammalian SNORF36 receptor additionally comprises an inducible promoter.

- 61. The transgenic, nonhuman mammal of claim 57 or 58, wherein the DNA encoding the mammalian SNORF36 receptor additionally comprises tissue specific regulatory elements.
- 62. A transgenic, nonhuman mammal of claim 57, 58, or 59, wherein the transgenic, nonhuman mammal is a mouse.
 - 63. A process for identifying a chemical compound which specifically binds to a mammalian SNORF36 receptor which comprises contacting cells containing DNA encoding, and expressing on their cell surface, the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with the compound under conditions suitable for binding, and detecting specific binding of the chemical compound to the mammalian SNORF36 receptor.
 - A process for identifying a chemical compound which 64. specifically binds to a mammalian SNORF36 receptor which comprises contacting a membrane preparation from cells containing DNA encoding, and expressing on their cell surface, the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with the compound under conditions suitable for binding, and detecting specific binding of chemical compound to the mammalian the receptor.
 - 65. The process of claim 63 or 64, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor.

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-140-

- 66. The process of claim 63 or 64, wherein the mammalian SNORF36 receptor has substantially the same amino acid sequence as the human SNORF36a receptor encoded by plasmid pcDNA3.1-hSNORF36a-f (ATCC Accession No. PTA-203977).
- 67. The process of claim 63 or 64, wherein the mammalian SNORF36 receptor has substantially the same amino acid sequence as the human SNORF36b receptor encoded by the plasmid pcDNA3.1-hSNORF36b-f (ATCC Accession No. PTA-203976).
- 68. The process of claim 63 or 64, wherein the mammalian SNORF36 receptor has substantially the same amino acid sequence as that shown in Figures 2A-2B (SEQ ID NO: 2) or Figures 4A-4C (SEQ ID NO: 4).
- 69. The process of claim 63 or 64, wherein the mammalian SNORF36 receptor has the amino acid sequence shown in Figures 2A-2B (SEQ ID NO: 2) or Figures 4A-4C (SEQ ID NO: 4).
 - 70. The process of claim 63 or 64, wherein the mammalian SNORF36 receptor is a rat SNORF36 receptor.
- 71. The process of claim 63 or 64, wherein the mammalian SNORF36 receptor has substantially the same amino acid sequence as the rat SNORF36 receptor encoded by plasmid pEXJ.T7-rSNORF36-f (ATCC Patent Depository No. PTA-1216).
 - 72. The process of claim 63 or 64, wherein the mammalian SNORF36 receptor has substantially the same amino acid sequence as that shown in Figures 10A-10C (SEQ ID NO: 8).

-141-

- 73. The process of claim 63 or 64, wherein the mammalian SNORF36 receptor has the same amino acid sequence as that shown in Figures 10A-10C (SEQ ID NO: 8).
- 5 74. The process of claim 63 or 64, wherein the compound is not previously known to bind to a mammalian SNORF36 receptor.
 - 75. A compound identified by the process of claim 74.
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- 76. A process of claim 63 or 64, wherein the cell is an insect cell.
- 77. The process of claim 63 or 64, wherein the cell is a mammalian cell.
 - 78. The process of claim 77, wherein the cell is nonneuronal in origin.
- 79. The process of claim 78, wherein the nonneuronal cell is a COS-7 cell, 293 human embryonic kidney cell, a CHO cell, a NIH-3T3 cell, a mouse Yl cell, or a LM(tk-) cell.
- 25 80. A process of claim 77, wherein the compound is a compound not previously known to bind to a mammalian SNORF36 receptor.
 - 81. A compound identified by the process of claim 80.
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82. A process involving competitive binding for identifying a chemical compound which specifically binds to a mammalian SNORF36 receptor which comprises separately contacting cells expressing on their cell surface the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with

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-142-

both the chemical compound and a second chemical compound known to bind to the receptor, and with only the second chemical compound, under conditions suitable for binding of such compounds to the receptor, and detecting specific binding of the chemical compound to the mammalian SNORF36 receptor, a decrease in the binding of the second chemical compound to the mammalian SNORF36 receptor in the presence of the chemical compound being tested indicating that such chemical compound binds to the mammalian SNORF36 receptor.

- A process involving competitive binding for identifying 83. a chemical compound which specifically binds to a mammalian SNORF36 receptor which comprises separately membrane preparation from cells a contacting expressing on their cell surface the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with both the chemical compound and a second chemical compound known to bind to the receptor, and with only the second chemical compound, under conditions suitable for binding of such compounds to the receptor, and detecting specific binding of the chemical compound to the mammalian SNORF36 receptor, a decrease in the binding of the second chemical compound to the mammalian SNORF36 receptor in the presence of the chemical compound being tested indicating that such chemical compound binds to the mammalian SNORF36 receptor.
- 84. A process of claim 82 or 83, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor.
- 85. A process of claim 82 or 83, wherein the mammalian SNORF36 receptor is a rat SNORF36 receptor.

-143-

- 86. The process of claim 82 or 83, wherein the cell is an insect cell.
- 87. The process of claim 82 or 83, wherein the cell is a mammalian cell.
- 88. The process of claim 87, wherein the cell is nonneuronal in origin.
- 10 89. The process of claim 88, wherein the nonneuronal cell is a COS-7 cell, 293 human embryonic kidney cell, a CHO cell, a NIH-3T3 cell, a mouse Yl cell, or a LM(tk-) cell.
- 15 90. The process of claim 89, wherein the compound is not previously known to bind to a mammalian SNORF36 receptor.
 - 91. A compound identified by the process of claim 90.

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92. A method of screening a plurality of chemical compounds not known to bind to a mammalian SNORF36 receptor to identify a compound which specifically binds to the mammalian SNORF36 receptor, which comprises

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(a) contacting cells transfected with, and expressing, DNA encoding the mammalian SNORF36 receptor with a compound known to bind specifically to the mammalian SNORF36 receptor;

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(b) contacting the cells of step (a) with the plurality of compounds not known to bind specifically to the mammalian SNORF36 receptor, under conditions permitting binding of compounds known to bind to the mammalian SNORF36 receptor;

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-144-

- (c) determining whether the binding of the compound known to bind to the mammalian SNORF36 receptor is reduced in the presence of the plurality of compounds, relative to the binding of the compound in the absence of the plurality of compounds; and if so
- (d) separately determining the binding to the mammalian SNORF36 receptor of each compound included in the plurality of compounds, so as to thereby identify any compound included therein which specifically binds to the mammalian SNORF36 receptor.
- 93. A method of screening a plurality of chemical compounds not known to bind to a mammalian SNORF36 receptor to identify a compound which specifically binds to the mammalian SNORF36 receptor, which comprises
 - (a) contacting a membrane preparation from cells transfected with, and expressing, DNA encoding the mammalian SNORF36 receptor with the plurality of compounds not known to bind specifically to the mammalian SNORF36 receptor under conditions permitting binding of compounds known to bind to the mammalian SNORF36 receptor;
 - (b) determining whether the binding of a compound known to bind to the mammalian SNORF36 receptor is reduced in the presence of the plurality of compounds, relative to the binding of the compound in the absence of the plurality of compounds; and if so
- 35 (c) separately determining the binding to the mammalian SNORF36 receptor of each compound

-145-

included in the plurality of compounds, so as to thereby identify any compound included therein which specifically binds to the mammalian SNORF36 receptor.

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- 94. A method of claim 92 or 93, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor or a human SNORF36b receptor.
- 95. A method of claim 92 or 93, wherein the mammalian SNORF36 receptor is a rat SNORF36 receptor.
 - 96. A method of claim 92 or 93, wherein the cell is a mammalian cell.

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- 97. A method of claim 96, wherein the mammalian cell is non-neuronal in origin.
- 98. The method of claim 97, wherein the non-neuronal cell is a COS-7 cell, a 293 human embryonic kidney cell, a LM(tk-) cell, a CHO cell, a mouse Y1 cell, or an NIH-3T3 cell.
- 99. A method of detecting expression of a mammalian SNORF36
 receptor by detecting the presence of mRNA coding for
 the mammalian SNORF36 receptor which comprises
 obtaining total mRNA from the cell and contacting the
 mRNA so obtained with the nucleic acid probe of claim
 36, 37, 38, 39, 40 or 41 under hybridizing conditions,
 detecting the presence of mRNA hybridized to the probe,
 and thereby detecting the expression of the mammalian
 SNORF36 receptor by the cell.
- 100. A method of detecting the presence of a mammalian SNORF36 receptor on the surface of a cell which comprises contacting the cell with the antibody of

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-146-

claim 47 under conditions permitting binding of the antibody to the receptor, detecting the presence of the antibody bound to the cell, and thereby detecting the presence of the mammalian SNORF36 receptor on the surface of the cell.

- 101. A method of determining the physiological effects of varying levels of activity of mammalian SNORF36 receptors which comprises producing a transgenic, nonhuman mammal of claim 57 whose levels of mammalian SNORF36 receptor activity are varied by use of an inducible promoter which regulates mammalian SNORF36 receptor expression.
- 102. A method of determining the physiological effects of varying levels of activity of mammalian SNORF36 receptors which comprises producing a panel of transgenic, nonhuman mammals of claim 57 each expressing a different amount of mammalian SNORF36 receptor.
 - 103. A method for identifying an antagonist capable of alleviating an abnormality wherein the abnormality is alleviated by decreasing the activity of a mammalian SNORF36 receptor comprising administering a compound to the transgenic, nonhuman mammal of claim 57, 58, or 59, and determining whether the compound alleviates any physiological and/or behavioral abnormality displayed by the transgenic, nonhuman mammal as a result of overactivity of a mammalian SNORF36 receptor, the alleviation of such an abnormality identifying the compound as an antagonist.
 - 104. The method of claim 103, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor, a human SNORF36b receptor or a rat SNORF36 receptor.

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-147-

- 105. An antagonist identified by the method of claim 103.
- 106. A composition comprising an antagonist of claim 105 and a carrier.
- 107. A method of treating an abnormality in a subject wherein the abnormality is alleviated by decreasing the activity of a mammalian SNORF36 receptor which comprises administering to the subject an effective amount of the pharmaceutical composition of claim 106, thereby treating the abnormality.
- for identifying an agonist capable of 108. A method alleviating an abnormality in a subject wherein the abnormality is alleviated by increasing the activity of 15 a mammalian SNORF36 receptor comprising administering a compound to the transgenic, nonhuman mammal of claim 57, 58, or 59, and determining whether the compound any physiological and/or behavioral alleviates abnormality displayed by the transgenic, nonhuman 20 mammal, the alleviation of such an abnormality identifying the compound as an agonist.
- 109. The method of claim 108, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor, a human SNORF36b receptor or a rat SNORF36 receptor.
 - 110. An agonist identified by the method of claim 109.
- 30 111. A composition comprising an agonist identified by the method of claim 110 and a carrier.
- 112. A method of treating an abnormality in a subject wherein the abnormality is alleviated by increasing the activity of a mammalian SNORF36 receptor which comprises administering to the subject an effective

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amount of the composition of claim 111 so as to thereby treat the abnormality.

- 113. A method for diagnosing a predisposition to a disorder associated with the activity of a specific mammalian allele which comprises:
 - (a) obtaining DNA of subjects suffering from the disorder;
 - (b) performing a restriction digest of the DNA with a panel of restriction enzymes;
 - (c) electrophoretically separating the resulting DNA fragments on a sizing gel;
 - (d) contacting the resulting gel with a nucleic acid probe capable of specifically hybridizing with a unique sequence included within the sequence of a nucleic acid molecule encoding a mammalian SNORF36 receptor and labeled with a detectable marker;
 - (e) detecting labeled bands which have hybridized to the DNA encoding a mammalian SNORF36 receptor of claim 1 to create a unique band pattern specific to the DNA of subjects suffering from the disorder;
 - (f) repeating steps (a)-(e) with DNA obtained for diagnosis from subjects not yet suffering from the disorder; and
 - (g) comparing the unique band pattern specific to the DNA of subjects suffering from the disorder from step (e) with the band pattern from step (f) for subjects not yet suffering from the disorder so as

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-149-

to determine whether the patterns are the same or different and thereby diagnose predisposition to the disorder if the patterns are the same.

- 5 114. The method of claim 113, wherein a disorder associated with the activity of a specific mammalian allele is diagnosed.
- 115. A method of preparing the purified mammalian SNORF36 receptor of claim 15 which comprises:
 - (a) culturing cells which express the mammalian SNORF36 receptor;
- 15 (b) recovering the mammalian SNORF36 receptor from the cells; and
 - (c) purifying the mammalian SNORF36 receptor so recovered.
 - 116. A method of preparing the purified mammalian SNORF36 receptor of claim 15 which comprises:
- (a) inserting a nucleic acid encoding the mammalian SNORF36 receptor into a suitable expression vector;
 - (b) introducing the resulting vector into a suitable host cell;
 - (c) placing the resulting host cell in suitable conditions permitting the production of the mammalian SNORF36 receptor;
- 35 (d) recovering the mammalian SNORF36 receptor so produced; and optionally

-150-

- (e) isolating and/or purifying the mammalian SNORF36 receptor so recovered.
- 117. A process for determining whether a chemical compound is a mammalian SNORF36 receptor agonist which comprises contacting cells transfected with and expressing DNA encoding the mammalian SNORF36 receptor with the compound under conditions permitting the activation of the mammalian SNORF36 receptor, and detecting any increase in mammalian SNORF36 receptor activity, so as to thereby determine whether the compound is a mammalian SNORF36 receptor agonist.

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- is a mammalian SNORF36 receptor antagonist which comprises contacting cells transfected with and expressing DNA encoding the mammalian SNORF36 receptor with the compound in the presence of a known mammalian SNORF36 receptor agonist, under conditions permitting the activation of the mammalian SNORF36 receptor, and detecting any decrease in mammalian SNORF36 receptor activity, so as to thereby determine whether the compound is a mammalian SNORF36 receptor antagonist.
- 25 119. A process of claim 117 or 118, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor, a human SNORF36b receptor or a rat SNORF36 receptor.
- 120. A composition which comprises an amount of a mammalian SNORF36 receptor agonist determined by the process of claim 117 effective to increase activity of a mammalian SNORF36 receptor and a carrier.
- 121. A composition of claim 120, wherein the mammalian SNORF36 receptor agonist is not previously known.

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-151-

122. A composition which comprises an amount of a mammalian SNORF36 receptor antagonist determined by the process of claim 118 effective to reduce activity of a mammalian SNORF36 receptor and a carrier.

123. A composition of claim 122, wherein the mammalian SNORF36 receptor antagonist is not previously known.

- 124. A process for determining whether a chemical compound specifically binds to and activates a mammalian SNORF36 10 receptor, which comprises contacting cells producing a second messenger response and expressing on their cell surface the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with the chemical compound under conditions 15 suitable for activation of the mammalian receptor, and measuring the second messenger response in the presence and in the absence of the chemical compound, a change in the second messenger response in the presence of the chemical compound indicating that 20 the compound activates the mammalian SNORF36 receptor.
 - 125. The process of claim 124, wherein the second messenger response comprises chloride channel activation and the change in second messenger is an increase in the level of chloride current.
 - 126. The process of claim 124, wherein the second messenger response comprises intracellular calcium levels and the change in second messenger is an increase in the measure of intracellular calcium.
- 127. The process of claim 124, wherein the second messenger response comprises release of inositol phosphate and the change in second messenger is an increase in the level of inositol phosphate.

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- 128. A process for determining whether a chemical compound specifically binds to and inhibits activation of a mammalian SNORF36 receptor, which comprises separately contacting cells producing a second messenger response and expressing on their cell surface the mammalian SNORF36 receptor, wherein such cells do not normally express the mammalian SNORF36 receptor, with both the chemical compound and a second chemical compound known to activate the mammalian SNORF36 receptor, and with only the second chemical compound, under conditions suitable for activation of the mammalian SNORF36 receptor, and measuring the second messenger response in the presence of only the second chemical compound and in the presence of both the second chemical compound and the chemical compound, a smaller change in the second messenger response in the presence of both the chemical compound and the second chemical compound than in the presence of only the second chemical compound indicating that the chemical compound inhibits activation of the mammalian SNORF36 receptor.
 - 129. The process of claim 128, wherein the second messenger response comprises chloride channel activation and the change in second messenger response is a smaller increase in the level of chloride current in the presence of both the chemical compound and the second chemical compound than in the presence of only the second chemical compound.
- 130. The process of claim 129, wherein the second messenger response comprises change in intracellular calcium levels and the change in second messenger response is a smaller increase in the measure of intracellular calcium in the presence of both the chemical compound and the second chemical compound than in the presence of only the second chemical compound.

-153-

- 131. The process of claim 128, wherein the second messenger response comprises release of inositol phosphate and the change in second messenger response is a smaller increase in the level of inositol phosphate in the presence of both the chemical compound and the second chemical compound than in the presence of only the second chemical compound.
- 132. A process of any of claims 124, 125, 126, 127, 128, 129, 130, or 131, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor, a human SNORF36b receptor or a rat SNORF36 receptor.
- 133. The process of any one of claims 121, 125, 126, 127, 128, 129, 130, or 131, wherein the cell is an insect cell.
- 134. The process of any one of claims 124, 125, 126, 127, 128, 129, 130, or 131, wherein the cell is a mammalian cell.
 - 135. The process of claim 134, wherein the mammalian cell is nonneuronal in origin.
- 25 136. The process of claim 135, wherein the nonneuronal cell is a COS-7 cell, CHO cell, 293 human embryonic kidney cell, NIH-3T3 cell or LM(tk-) cell.
- or 131, wherein the compound is not previously known to bind to a mammalian SNORF36 receptor.
 - 138. A compound determined by the process of claim 137.
- 35 139. A composition which comprises an amount of a mammalian SNORF36 receptor agonist determined to be such by the

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-154-

process of claim 124, 125, 126, or 127, effective to increase activity of the mammalian SNORF36 receptor and a carrier.

- 5 140. A composition of claim 139, wherein the mammalian SNORF36 receptor agonist is not previously known.
 - 141. A composition which comprises an amount of a mammalian SNORF36 receptor antagonist determined to be such by the process of claim 128, 129, 130, or 131, effective to reduce activity of the mammalian SNORF36 receptor and a carrier.
- 142. A composition of claim 141, wherein the mammalian SNORF36 receptor antagonist is not previously known.
 - 143. A method of screening a plurality of chemical compounds not known to activate a mammalian SNORF36 receptor to identify a compound which activates the mammalian SNORF36 receptor which comprises:
 - (a) contacting cells transfected with and expressing the mammalian SNORF36 receptor with the plurality of compounds not known to activate the mammalian SNORF36 receptor, under conditions permitting activation of the mammalian SNORF36 receptor;
 - (b) determining whether the activity of the mammalian SNORF36 receptor is increased in the presence of one or more of the compounds; and if so
 - (c) separately determining whether the activation of the mammalian SNORF36 receptor is increased by any compound included in the plurality of compounds, so as to thereby identify each compound which activates the mammalian SNORF36 receptor.

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-155-

- 144. A method of claim 143, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor, a human SNORF36b receptor or a rat SNORF36 receptor.
- 145. A method of screening a plurality of chemical compounds not known to inhibit the activation of a mammalian SNORF36 receptor to identify a compound which inhibits the activation of the mammalian SNORF36 receptor, which comprises:

(a) contacting cells transfected with and expressing the mammalian SNORF36 receptor with the plurality of compounds in the presence of a known mammalian SNORF36 receptor agonist, under conditions permitting activation of the mammalian SNORF36 receptor;

- (b) determining whether the extent or amount of activation of the mammalian SNORF36 receptor is reduced in the presence of one or more of the compounds, relative to the extent or amount of activation of the mammalian SNORF36 receptor in the absence of such one or more compounds; and if so
- (c) separately determining whether each such compound inhibits activation of the mammalian SNORF36 receptor for each compound included in the plurality of compounds, so as to thereby identify any compound included in such plurality of compounds which inhibits the activation of the mammalian SNORF36 receptor.
- 146. A method of claim 145, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor, a human SNORF36b receptor or rat SNORF36 receptor.

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-156-

- 147. A method of any one of claims 143, 144, 145 or 146, wherein the cell is a mammalian cell.
- 148. A method of claim 147, wherein the mammalian cell is non-neuronal in origin.
- 149. The method of claim 148, wherein the non-neuronal cell is a COS-7 cell, a 293 human embryonic kidney cell, a LM(tk-) cell or an NIH-3T3 cell.
- 150. A composition comprising a compound identified by the method of claim 143 or 144 in an amount effective to increase mammalian SNORF36 receptor activity and a carrier.
- 151. A composition comprising a compound identified by the method of claim 145 or 146 in an amount effective to decrease mammalian SNORF36 receptor activity and a carrier.
 - 152. A method of treating an abnormality in a subject wherein the abnormality is alleviated by increasing the activity of a mammalian SNORF36 receptor which comprises administering to the subject a compound which is a mammalian SNORF36 receptor agonist in an amount effective to treat the abnormality.
 - 153. A method of treating an abnormality in a subject wherein the abnormality is alleviated by decreasing the activity of a mammalian SNORF36 receptor which comprises administering to the subject a compound which is a mammalian SNORF36 receptor antagonist in an amount effective to treat the abnormality.
- 35 154. A process for making a composition of matter which specifically binds to a mammalian SNORF36 receptor

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-157-

which comprises identifying a chemical compound using the process of any one of claims 63, 64, 82, 83, 92 or 93 and then synthesizing the chemical compound or a novel structural and functional analog or homolog thereof.

- 155. The process of claims 154, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor, a human SNORF36b receptor or a rat SNORF36 receptor.
- 156. A process for making a composition of matter which specifically binds to a mammalian SNORF36 receptor which comprises identifying a chemical compound using the process of any of claims 117, 124, or 143 and then synthesizing the chemical compound or a novel structural and functional analog or homolog thereof.
 - 157. The process of claim 156, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor, a human SNORF36b receptor or a rat SNORF36 receptor.
 - 158. A process for making a composition of matter which specifically binds to a mammalian SNORF36 receptor which comprises identifying a chemical compound using the process of any of claims 118, 128 or 145 and then synthesizing the chemical compound or a novel structural and functional analog or homolog thereof.
- 159. The process of claim 158, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor, a human SNORF36b receptor or a rat SNORF36 receptor.
 - 160. A process for preparing a composition which comprises admixing a carrier and a pharmaceutically effective amount of a chemical compound identified by the process of any of claims 63, 64, 82, 83, 92 or 93 or a novel

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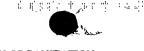
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structural and functional analog or homolog thereof.

- 161. The process of claim 160, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor, a human SNORF36b receptor or a rat SNORF36 receptor.
- 162. A process for preparing a composition which comprises admixing a carrier and a pharmaceutically effective amount of a chemical compound identified by the process of any of claims 117, 124, or 143 or a novel structural and functional analog or homolog thereof.
 - 163. The process of claim 162, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor, a human SNORF36b receptor or a rat SNORF36 receptor.
 - 164. A process for preparing a composition which comprises admixing a carrier and a pharmaceutically effective amount of a chemical compound identified by the process of any of claims 118, 128 or 145 or a novel structural and functional analog or homolog thereof.
- 165. The process of claim 164, wherein the mammalian SNORF36 receptor is a human SNORF36a receptor, a human SNORF36b receptor or a rat SNORF36 receptor.







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(54) Title: DNA ENCODING SNORF36a AND SNORF36b RECEPTORS

(57) Abstract

This invention provides isolated nucleic acids encoding mammalian SNORF36 receptors, purified mammalian SNORF36 receptors, vectors comprising nucleic acid encoding mammalian SNORF36 receptors, cells comprising such vectors, antibodies directed to mammalian SNORF36 receptors, nucleic acid probes useful for detecting nucleic acid encoding mammalian SNORF36 receptors, antisense oligonucleotides complementary to unique sequences of nucleic acid encoding mammalian SNORF36 receptors, transgenic, nonhuman animals which express DNA encoding normal or mutant mammalian SNORF36 receptors, methods of isolating mammalian SNORF36 receptors, methods of treating an abnormality that is linked to the activity of the mammalian SNORF36 receptors, as well as methods of determining binding of compounds to mammalian SNORF36 receptors, methods of identifying agonists and antagonists of SNORF36 receptors, and agonists and antagonists so identified.

WO 00/66630

FIGURE 1A

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	CAACTCAGG <u>ATG</u> AACCCTCCTTCGGGGCCAAGAGTCCCGCCCAGCCCA	09	
61	AGCTGCATGCCACCCCAGCACCACCCAGCTGGGACAGCTCCCAGAGCAGCATCTCC	1.20	
121	AGUCTIGGGCCCGGCTTCCATCCATCCCACACCACCTGGGACTTGGGCTGCTGCCTGG	180	
181	GTCCCCCCCCCACGGTTGALGTTCCAGACCATGCCCACTATACCCTGGGCACAGTGATC	240	
241	TTGCTGGTGGGACTCACGGGGATGCTGGGCAACCTGACGGTCATCTATACCTTCTGCAGG	300	1/
301	AGCAGAAGCCTCCGGACACCTGCCAACATGTTCATTATCAACCTCGCGGTCAGCGACTTC	360	33
361	CTCATGTCCTTCACCCAGGCCCCTGTCTTCTTCACCAGTAGCCTCTATAAGCAGTGGCTC	420	
421	TTTGGGGAGACAGGCTGCGAGTTCTATGCCTTCTGTGGAGCTCTCTTTGGCATTTTCCTCC	480	
481	ATGATCACCCTGACGCCCATCGCCCTGGACCGCTACCTGGTAATCACACGCCCGCTGGCC	540	
541	ACCTTTGGTGTGCGTCCAAGAGGCGTGCGGCATTTGTCCTGCTGGGCGTTTGGCTCTAT	009	
601	GCCCTGGCCTGGAGTCTGCCACCCTTCTTCGGCTGGAGCGCCTACGTGCCCCGAGGGGTTG	099	
661	CTGACATCCTGCTCCTGGGACTACATGAGCTTCACGCCGGCCG	720	

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FIGURE 1B

7	721	CTTCTCTGTGCTTCGTGTTCTTCCTCCTCTGCTTATCATCATCTACTGCTACATCTTC	780	
7	781	ATCTTCAGGGCCATCCGGGAGACAGGACGGGCTCTCCAGACCTTCGGGGCCTGCAAGGGC	840	-
ω SU	841	. AATGGCGAGTCCCTGTGGCAGCGGCAGCGGCTGCAGAGTGCAAGATGGCCAAGATC	900	
O) RSTITI	901	ATGCTGCTGGTCATCCTCCTCTTCGTGCTCTCTGGGCTCCCTATTCCGCTGTGGCCCTG	960	
の UTF SH	961	GTGGCCTTTGCTGGGTACGCACGTCCTGACACCCTACATGAGCTCGGTGCCAGCCGTC	1020	2/
인 IEET (E	1021	ATCGCCAAGGCCTCTGCAATCCACAACCCCATTATTACGCCATCACCCCACCCA	1080	'3 5
0 UI F 2	1081	AGGGTGGCCATTGCCCAGCACCTGCCCTGGGGGTGTGCTGGGTGTATCACGCCGG	1140	
	1141	CACAGTCGCCCCTACCCCTACCGCTCCACCCACCGCTCCACGCTGACCAGCCACCC	1200	
12	1201	TCCAACCTCAGCTGGATCTCCATACGGAGGCGCCAGGAGTCCCTGGGCTCGGAGAGTGAG	1260	
12	1261	GTGGGCTGGACACACATGGAGGCAGCTGTTGTGGGGAGCTGCCCAGCAAGCA	1320	
13	1321	CGGTCCCTCTACGGTCAGGGTCTGGAGGACTTGGAAGCCAAGGCACCCCCCAGACCCCAG	1380	
13	1381	GGACACGAAGCAGAGACTCCAGGGAAGACCAAGGGGCTGATCCCCCAGCCAG	1440	

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FIGURE 1C

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FIGURE 31

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	099	GCGGCATTTGTCCTGCTGGCGTTTGGCTCTATGCCCTGGCCTGGAGTCTGCCACCCTTC
	009	GACCGCTACCTGGTAATCACACGCCCGCTGGCCACCTTTGGTGTGGCGTCCAAGAGGCGT
-	540	GCCTTCTGGAGCTCTCTTTGGCATTTCCTCCATGATCACCCCTGACGGCCATCGCCCTG
	480	TTCȚTCACCAGTAGCCTCTATAAGCAGTGGCTCTTTGGGGAGACAGGCTGCGAGTTCTAT
	420	ATGTTCATTATCAACCTCGCGGTCAGCGACTTCCTCATGTCCTTCACCCCAGGCCCCTGTC
′ 35	360	GCTGTGTGCTTCGTGGAGTCACTGTGATGATGCAGAGCCAGAAGCCTCCGGACACCTGCCAAC
6/	300	TTGCTGGTGGGACTCACGGGGATGCTGGGCAACCTGACGGTCATCTATACCTTCTGCAGA
	240	GTCCCCCTCCCCACGGTTGATGTTCCAGACCATGCCCACTATACCCTGGGCACAGTGATC
	180	AGCCTGGGCCGGCTTCCATCCATCCACACCACACCTGGGACTTGGGCTGCTGCTGG
	120	AGCTGC <u>ATG</u> GCCACCCCAGCACCCAGCTGGTGGGACAGCTCCCAGAGCAGCATCTCC
	09	CAACTCAGGATGAACCCTCCTTCGGGGCCAAGAGTCCCGCCCAGCCCAACCCAAGAGCCC

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FIGURE 31

780	840	006	096	7/020	35 0801	1140	1200	1260	1320	1380	1440
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1 AGCTTCACGCCGGCCGTGCCTACACCCATGCTTCTCTGCTTCGTGTTCTTCCTC	1 CCTCTGCTTATCATCTACTGCTACATCTTCATCTTCAGGGCCATCCGGGAGACAGGA	1 CGGGCTCTCCAGACCTTCGGGGCCTGCAAGGGCAATGGCGAGTCCCTGTGGCAGCGGCAG	1 CGGCTGCAGAGGTGCAAGATGGCCAAGATCATGCTGCTGGTCATCCTCCTCTTCGTG	1 CTCTCCTGGGCTCCCTATTCCGCTGTGGCCCTTGGCCTTTGCTGGGTACGCACGTC	1 CTGACACCCTACATGAGCTCGGTGCCAGCCGTCATCGCCAAGGCCTCTGCAATCCACAAC	1 CCCATCATTTACGCCATCACCCCCAAGTACAGGGTGGCCATTGCCCAGCACCTGCCC	1 TGCCTGGGGGTGTGTATCACGCCGGCACAGTCGCCCCTACCCCAGCTACCGC	1 TCCACCCACGCTCCACGCTGACCAGCCACCTCCAACCTCAGCTGGATCTCCATACGG	1 AGGCGCCAGGAGTCCCTGGGCTCGGAGAGTGAGGTGGGCTGGACACACATGGAGGCAGCA	1 GCTGTGTGGGGAGCTGCCCAGCAAGCAAATGGGCGGTCCCTCTACGGTCAGGGTCTGGAG	1 GACTTGGAAGCCAAGGCACCCCCAGACCCCAGGGACACGAAGCAGAGACTCCAGGGAAG
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FIGURE3C

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ACCAAGGGGCTGATCCCCAGCCAGGACCCCAGGATGTAGGACGCCCACTGGCTCTCCCTT TCTTCTGAGACACATCCAGCCCCCCCCCCCCCTCATAT 1501 1441

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FIGURE 4A

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FIGURE 4B

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260	280	300	320	340	360	380	400	420	440	460	480
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241	261	281	301	321	341	361	381	401	421	441	461

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FIGURE 4C

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09	120	180	240	250	
. CATAGCCATGGACCGCTATCTGGTGATCACACGTCCACTGGCCACCATCGGCATGAGATC	. CAAGAGACGGACGGCACTAGTCCTGCTAGGTGTCTGGCTCTATGCCCTGGCCTGGAGTCT	GCCGCCTTTCTTTGGCTGGAGCGCCTACGTGCCCGAGGGGGCTGCTGACATCCTGCTCCTG	GGACTACGTGACCTTCACGCCCCTCGTGCGCGCCTACACCATGCTGCTCTTCTGCTTTTGT	LCTTCTTCCTC	
Π	61	121	181	241	
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FIGURE 71

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212	250
GCTGACATCCTGCTCCTGGGACTACGTGACCTTCACGCCCCTCGTGCGCGGGGGIIIIIIIIII	CCTACACCATGCTGCTTTCTTGTCTTCTTCCTC
163 651	213
Rat SNORF36 Hum SNORF36	Rat SNORF36 Hum SNORF36
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FIGURE

250 IAMDRYLVITRPLATIGMRSKRRTALVLLGVWLYALA 201 Hum SNORF36

Hum SNORF36

Hum SNORF36 SNORF36 Rat

FIGURE 9A

			1	7/35	5					
09	120	180	240	300	360	420	480	540	009	099
	TTAACTCAGGATCCCAGCTTTACCGCCAGCCCTGCCCTCCTACAAGGCATTTGGAACAGC	ACTCAGAACATCTCCGTCAGAGTCCAGCTTCTATCCGTTAGCCCCCACGACACCTGGGCTT	CAGGCTGCTGCCTGCCCTTCCCCACAGTCGACGTCCCAGATCATGCTCACTATACC	CTAGGCACGGTGATCCTGCTGGGGACTCACAGGGATGCTGGGTAACCTGACAGTCATC	TACACCTTCTGCAGGAATAGAGGCCTGCGGACACCGGCAAACATGCTCATCATCAACCTG	GCAGTCAGCGACTTCCTTATGTCGTTCACTCAGGCCCCGGTCTTCTTTGCCAGCAGCCTC	TACAAGAAGTGGCTCTTCGGGGAGACAGGTTGCAAGTTCTATGCCTTCTGTGGGGCTGTC	TTTGGCATCGTTTCCATGATCACCCTGACAGCCATAGCCATGGACCGCTATCTGGTGATC	ACACGTCCACTGGCCACCATCGGCATGAGATCCAAGAGACGGACG	GGTGTCTGGCTCTATGCCCTGGCCTGGAGTCTGCCGCCTTTCTTT
	61	177 173 178 178 178 178 178 178 178 178 178 178	181	241	301	361	421	481	541	601

FIGURE 9B

	1320	1261 AGTGAAGTGGGCTGGACAGACACAGAACAACAGCTGCGTGGGGAGCTGCCCAGCAAGCA
	1260	1201 CAGTCCTCAGACCTCAGCTGGATCTCTGGGCAGAAGCGCCAAGAGTCCCTGGGTTCTGAG
	1200	1141 GGCCAGCCACCCCTCCTCAGCTACCGCTCTACCCATCGCTCCACACTGAGCAGC
	1140	1081 AAGTACAGGGGGCGTTTGCTCAGCACTTGCCTTGGGGTGCTTCTTGGAGTATCA
	1080	1021 GCCGTCATTGCCAAGGCCTCGGCCATCCACAATCCTATCATCTATGCCATCACTCAC
5	1020	961 GCCCTGGTGGGCTTTGCTGGGTACTCGCACATCCTGACGCCCTACATGAGCTCGGTGCCA
8/3	960	901 AAGGTCGCACTGATCGTCATTCTCTCTTTGTGCTGTCCTGGGCTCCCTACTCCACTGTG
1	006	841 GGTGAGTCCCCTCTGCGGCGGCGGCAGTGGCAGCGGCTACAGAGTGAATGGAAGATGGCC
	840	TTCTGCTACATCTTCATCTTCAGGGCCATTCGAGAGACAGGCCGGGCCTGTGAGGGCTGT
-	780	CGCGCCTACACCATGCTCTTCTTCTTCTTCTTCCTCCCTC
	720	GTGCCCGAGGGGCTGCTGACATCCTGCTCCTGGGACTACGTGACCTTCACGCCCCTCGTG

FIGURE 9C

19/35

1321	AGTGGACAATCCTTCTGCAGTCATGACCTGGAAGATGGAGAGTCAAGGCTCCTTCCAGC	138
1381	CCCCAGGAACAGAATCCAAGACTCCCAAGACCAAGAGACACCTCCCCAGTCTGGACCGA	144
1441	AGGATGTAGGATGCCCAGTCCCCTCCCT	147

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FIGURE 10A

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FIGURE 11

400 300 350 200 200 250 250 300 150 150 CAGCTGCATGGCCACCCCAGCACCCAGCTGGTGGGACAGCTCCCAGA 100 CAGCTTTACCGCCAGCCCTGCCTCCTACAAGGCATTTGGAACAGCACTC 100 50 GGATGCTGGGTAACCTGACAGTCATCTACACCTTCTGCAGGAATAGAGGC AGAACATCTCCGTCAGAGTCCAGCTTCTATCCGTTAGCCCCACGACACCT GCAGCATCTCCAGCCTGGGCCGGCTTCCATCCATCAGTCCCACAGGCACCT ATGAACTCTCCTTCAGAATCAAGAGTCCCTTCAAGCTTAACTCAGGATCC 101 251 301 351 101 201 301 351 151 151 201 251 \leftarrow 51 21 hSNORF36a hSNORF36a hSNORF36a hSNORF36a hSNORF36a hSNORF36a hSNORF36a hSNORF36a rSNORF36 rSNORF36 rSNORF36 rSNORF36 rSNORF36 rSNORF36 rSNORF36 rSNORF36

24/35

hSNORF36a	401 AGCZ		450
rSNORF36	401 AGA	AGAAGTGGCTCTTCGGGGAGACAGGTTGCAAGTTCTATGCCTTCTGTGGG	450
hSNORF36a	451 GCT(200
rSNORF36	451 GCT	GCTGTCTTTGGCATCGTTTCCATGATCACCCTGACAGCCATAGCCATGGA	200
hSNORF36a	501 CCG		550
rSNORF36	501 CCG	CCGCTATCTGGTGATCACGTCCACTGGCCACCATCGGCATGAGATCCA	550
hSNORF36a	551 AGA(009
rSNORF36	551 AGA	AGAGACGGACTAGTCCTGCTAGGTGTCTGGCTCTATGCCCTGGCC	009
hSNORF36a	601 TGG	TGGAGTCTGCCACCCTTCTTCGGCTGGAGCGCCTACGTGCCCGAGGGGTT	650
rSNORF36	601 TGG	AGTCTGCCGCCTTTCTTTGGCTGGAGCGCCTACGTGCCCGAGGGGCT	650
hSNORF36a	651 GCT		700
rSNORF36	651 GCT	GCTGACATCCTGCTCCTGGGACTACGTGACCTTCACGCCCCTCGTGCGCG	700
hSNORF36a	701 CCT	CCTACACCATGCTTCTCTGCTTCGTGTTCTTCCTCCCTCTGCTTATC	750
rSNORF36	701 CCT	ACACCATGCTGCTCTTCTGCTTTGTCTTCCTCCTCTCTGCTCATT	750
hSNORF36a	751 ATC	ATCATCTACTGCTACATCTTCATCTTCAGGGCCATCCGGGAGACAGGACG	800
rSNORF36	751 ATC		800

FIGURE 11C

hSNORF36a	801	801 GGCTCTCCAGACCTTCGGGGCCTGCAAGGGCAATGGCGAGTCCCTGTGGC 850
rSNORF36	801	
hSNORF36a	851	AGCGGCAGCGGCTGCAGAGCGAGTGCCAAGATGGCCAAGATCATGCTGCTG 900
rSNORF36	842	
hSNORF36a	901	
rSNORF36	892	GTCATTCTCCTCTTTGTGCTGTCCTGGGCTCCCTACTCCACTGTGGCCCT 941
hSNORF36a	951	
rSNORF36	942	GGTGGGCTTTGCTGGGTACTCGCACATCCTGACGCCCTACATGAGCTCGG 991
hSNORF36a	1001	TGCCAGCCGTCATCGCCAAGGCCTCTGCAATCCACAACCCCATCATTTAC 1050
rSNORF36	992	TGCCAGCCGTCATTGCCAAGGCCTCGGCCATCCACAATCCTATCATCTAT 1041
hSNORF36a	1051	
rSNORF36	1042	GCCATCACTCACCCCAAGTACAGGGCGGCCATTGCTCAGCACTTGCCTTG 1091
hSNORF36a	1101	CCTGGGGGTGCTGGGTGTATCACGCCGGCACAGTCGCCCCTACCCCA 1150
rSNORF36	1092	
hSNORF36a	1151	GCTACCGCTCCACCCACCGCTCCACGCTGACCACCACCTCCAACCTC 1200
rSNORF36	1142	1142 GCTACCGCTCTACCCATCGCTCCACACTGAGCAGCCAGTCCTCAGACCTC 1191

FIGURE 11D

1SNORF 36a	1201	1SNORF36a 1201 AGCTGGATCTCCATACGGAGGCGCCAGGAGTCCCTGGGCTCGGAGTGA 1250	1250
rSNORF36	1192	-U	1241
SNORF36a	1251	1251 GGTGGGCTGGACACACATGGAGGCAGCTGTGTGGGGAGCTGCCCAGC 1300	1300
cSNORF36	1242	1242 AGTGGGCTGGACAGACAGAACAACAGCTGCGTGGGGAGCTGCCGAGC 1291	1291
nSNORF36a	1301	1301 AAGCAAATGGGCGGTCCCTCTACGGTCAGGGTCTGGAGGACTTGGAAGCC 1350	1350
rSNORF36	1292	1292 AAGCAAGTGGACAATCCTTCTGCAGTCATGACCTGGAAGATGGAGAAGTC 1341	1341
hSNORF36a	1351	1351 AAGGCACCCCCAGACCCCAGGGACACGAAGCAGAGACTCCAGGGAAGAC 1400	1400
rSNORF36	1342	1342 AAGGCTCCTTCCAGCCCCCCAGGAACAGAATCCAAGACTCCCAAGAC	1388
hSNORF36a	1401	1401 CAAGGGGCTGATCCCCAGGACCCCAGGATGTAG	1437
rSNORF36	1389		1425

1 MNPPSGPRVPPSPTQEPSCMATPA.PPSWWDSSQSSISSLGRLPSISPTA
GTWAA
LRTPANI
GLRTPANMLIINLAVSDFLMSFTQAPVFFASSLYKKWLFGETGCKFYAFC
SALFGISS
GAVFGIVSMITLTAIAMDRYLVITRPLATIGMRSKRRTALVLLGVWLYAL
AWSLPPFFGWSAYVPEGLLTSCSWDYMSFTPAVRAYTMLLCCFVFFLPLL
WSLPPFF
IIIYCYIFIFRAIRETGRALQTFGACKGNGESLWQRQRLQSECKMAK
IIFCYIF
MLLVILL
OALIVILLFVLSWAPYSTVALVGFAGYSHILTPYMSSVPAVIAKASAIHN
PIIYAITHPKYRVAIAQHLPCLGVLLGVSRRHSRPYPSYRSTHRSTLTSH
IIYAITH

FIGURE 12B

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28/35

TSNLSWISIRRRQESLGSESEVGWTHMEAAAVWGAAQQANGRSLYGQGLE 446 hSNORF36a hSNORF36a rSNORF36 rSNORF36

WO 00/66630

29/35

FIGURE 13A

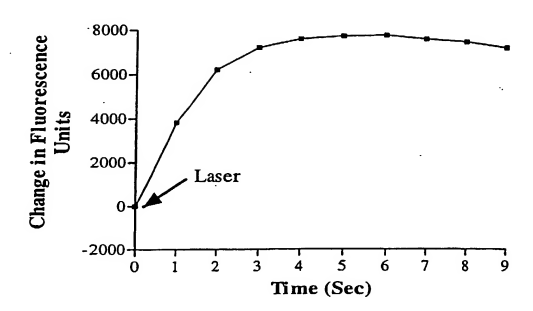
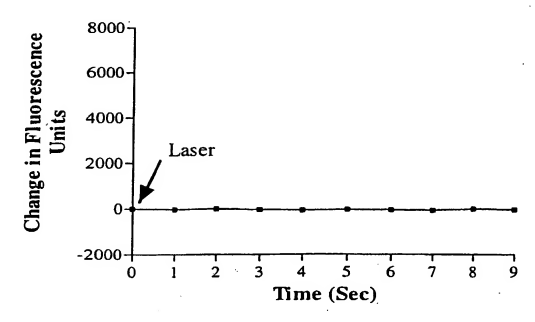


FIGURE 13B



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30/35

FIGURE 14A

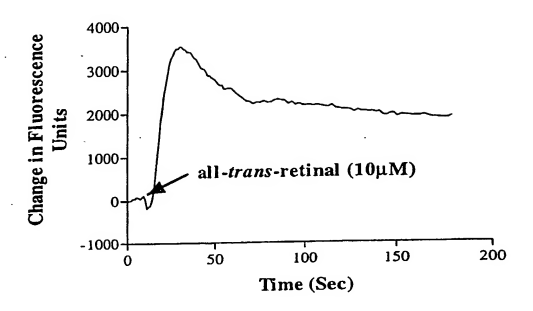
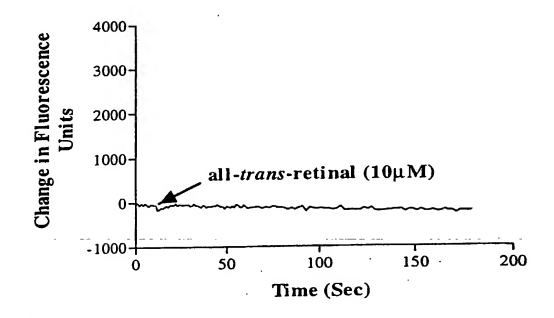


FIGURE 14B



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31/35

FIGURE 15A

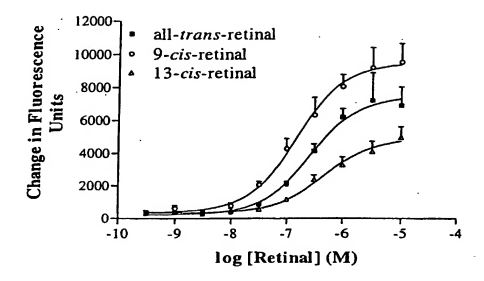
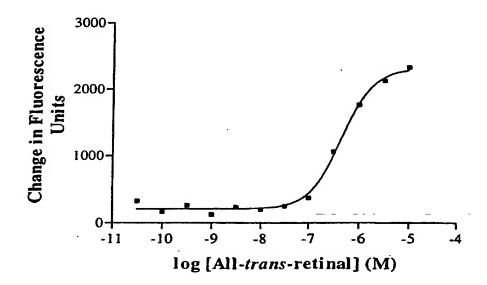


FIGURE 15B



32/35

FIGURE 16A

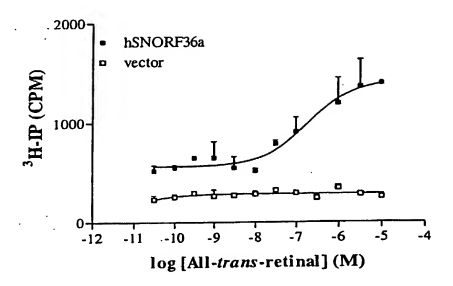
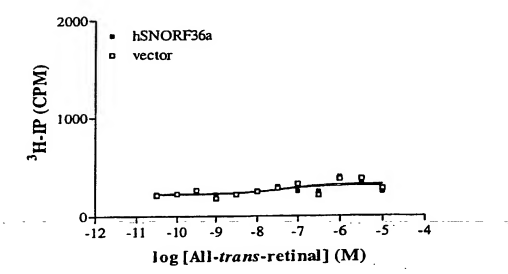


FIGURE 16B



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FIGURE 17A

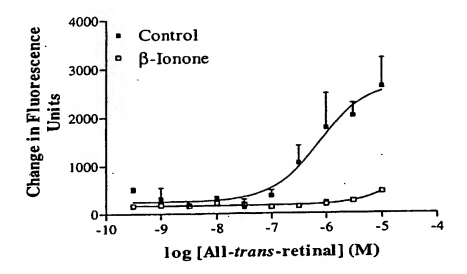
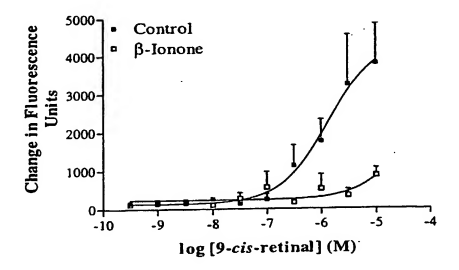


FIGURE 17B



34/35

FIGURE 18A



FIGURE 18B

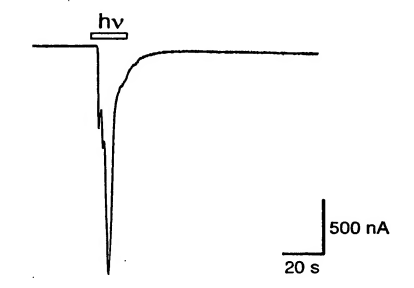
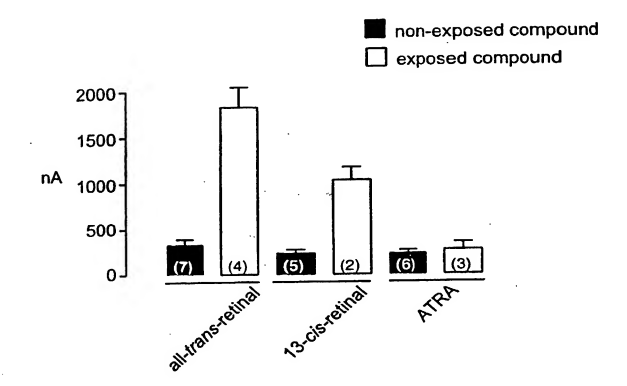


FIGURE 18C

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Figure 19



1795/59138-B-PCT-US/JPW/FHB

DECLARATION AND POWER OF ATTORNEY

As a below-named inventor, I hereby declare that:

My residence, post office address, and citizenship are as stated below next to my name.

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

the specification of	(which:				
	is allached h	nereio.			
•	x was filed on	November 1,	2001		_ as
	Application Serial No	<u>, 10/018,192</u>			 -
٠.	and was amended	November 1,	2001		
				(if applicable)	

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose to the U.S. Patent and Trademark Office all information known to me to be material to patentability as defined in Title 37, Code of Federal Regulations, Section 1.56.

I hereby claim foreign priority benefits under Title 35. United States Code, Section 119 (a)-(d) or Section 365(b) of any foreign application(s) for patent or inventor's certificate, or Section 365(a) of any PCT International Application which designated at least one country other than the United States, listed below. I have also identified below any foreign application for patent or inventor's certificate, or PCT International Application having a filing date before that of the earliest application from which priority is claimed:

Prior Foreign App	lication(s)		Priority Claimed			
Number	Country	Filing Date	<u>Yes</u>	<u>No</u>		
PCT/US00/12065	PCT	3 May 2000	<u>X</u>			
				-		
				-		
			- U.S. Serial N	Beth E. Borowsky et al. Io.: 10/018,192		
			Entered U.S.	National Stage:		

November 1, 2001 Exhibit B

Declaration and Power of Attorney		Page ?
I hereby claim the benefit under I provisional application(s) listed belo	iile 35. Uniied Siaies Code. Se ow:	ction 119(e) of any United State.
Provisional Application No.	Filing Dave	<u>Storus</u>
N/A	<u> </u>	
·		

I hereby claim the benefit under Title 35. United States Code. Section 120 of any United States Application(s), or Section 365(c) of any PCT International Application(s) designating the United States listed below. Insofar as this application discloses and claims subject matter in addition to that disclosed in any such prior Application in the manner provided by the first paragraph of Title 35. United States Code. Section 112. I acknowledge the duty to disclose to the United States Patent and Trademark Office all information known to me to be material to patentability as defined in Title 37. Code of Federal Regulations, Section 1.56. which became available between the filing date(s) of such prior Application(s), and the national or PCT international filing date of this application:

<u>Application Serial No.</u>	Filing Dois	Stones .
09/518,914	3 March 2000	Pending
09/303,593	3 May 1999	Abandoned
PCT/US00/12065	3 May 2000 ·	Pending

And I hereby oppoint

John P. White (Reg. No. 28,678); Christopher C. Dunham (Reg. No. 22,031); Norman H. Zivin (Reg. No. 25,385); Jay H. Maioli (Reg. No. 27,213); William E. Pelton (Reg. No. 25,702); Robert D. Katz (Reg. No. 30,141); Peter J. Philips (Reg. No. 29,691); Wendy E. Miller (Reg. No. 35,615); Richard S. Milner (Reg. No. 33,970); Roberto T. Maldonado (Reg. No. 38,232); Paul Teng (Reg. No. 40,837); Richard F. Jaworski (Reg. No 33,515); Pedro C. Fernandez (Reg. No. 41,741); Gary J. Gershik (Reg. No. 39,992); Spencer H. Schneider (Reg. No. 45,923); Alan J. Morrison (Reg. No. 37,399); Alan D. Miller (Reg. No. 42,889); and Frank Bruno (Reg. No. 46,583)

and each of them, all c/o Cooper & Dunham LLP. 1185 Avenue of the Americas, New York. New York 10036, my ottorneys, each with full power of substitution and revocation, to prosecute this application, to make alterations and amendments therein, to receive the patent, to transact all business in the Patent and Trademark Office connected therewith and to file any International Applications which are based thereon under the provisions of the Potent Cooperation Treaty.

Page 3

	Please address all communications, and direct all telephone calls, regarding this application to:
	John P. White
	Cooper & Dunham LLP
	1185 Avenue of the Americas New York, New York 10036
	Tel. (212) 278-0400
	I hereby declare that all statements made herein of my own knowledge are true and that all statem made on information and belief are believed to be true: and further that these statements were made the knowledge that willful false statements and the like so made are punishable by fine or imprisonm or both, under Section 1001 of Title 18 of the United States Code and that such wiilful false statem may jeopardize the validity of the application or any patent issued thereon.
	Full name of sole or first joint inventorBeth E. Borowsky
	Total Community of the
	Inventor's signature / Sulf forces
	Citizenship United States of America Date of signature 11/5/01
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)	Full name of joint Inventor (if any) Kristine L. Ogozalek
	V 1 1 () = 1 A
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	Full name of joint
	invenior (if ony) Parul P. Lakhlani
	Inventor's signature
	Inventor's signature
(Citizenship India Date of signature

Declaration and Power of Attorney

Page 3

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	Full name of sole or			
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	Inventor's signature			_
	Citizenship United States of America Da	ie of signature_		
	Residence 218 Park Street, Montclair,			
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	Full name of joint inventor (if any) Kristine L. Ogozalek		<u>.</u>	-
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